

Principles and Practice of

GERIATRIC

PSYCHIATRY

Second Edition

Edited by

John R. M. Copeland

Mohammed T. Abou-Saleh

Dan G. Blazer

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Preface

The editors were very gratified that the first edition of this textbook was generally well received and that a second edition has been called for. It is now seven years since the original book appeared, and there have been many more advances in the subject. In spite of new sections and some wholesale rewriting, it has been possible once again, to contain the information in one volume. Very sadly some of our original contributors have died. New authors have replaced them while others have been added in an endeavour to keep the text authoritative and up-to-date. The helpful criticisms of the first edition have been carefully considered in the

preparation of this one. Having so many distinguished authors with such a breadth of interest, while greatly enhancing the book, has led to a long gestation period, but we believe that it has been worthwhile. Much of the original format has been retained in order to continue to stimulate lively debate and exchange of views. If the book contributes to the growing strength of Geriatric Psychiatry internationally, it will have done its work.

John R. M. Copeland
Mohammed T. Abou-Saleh
Dan G. Blazer

Preface to First Edition

The discipline of the psychiatry of old age has moved rapidly in recent years and the number of practitioners has expanded world-wide. An authoritative text is required which draws on the knowledge of these experts and which reflects both new scientific advances and innovations in service development.

In a comparatively new subject many of the issues are still contentious and on some of these we have tried to provide the opportunity for the expression of different points of view. Readers are asked to judge the issues for themselves from the evidence set out.

Here and there short, special articles have been commissioned which present research findings in more detail and describe new aspects of care. They are intended to enliven the text and their choice has been dependent on timing and opportunity.

We have also tried to give a “feel” for what is happening in developing countries and the scope of the problems experienced by local practitioners.

Even a book of this size can never be complete and no doubt gaps in the coverage of subjects will be identified. We would be glad to have them pointed out. The more comprehensive a book aims to be the longer it takes to come to publication and in a fast-moving area of knowledge this can be a problem. Many of our authors have been kind enough to update their contributions at a late stage, which we hope has overcome this difficulty to some extent.

In the early stages of the development of a subject there is insufficient corpus of knowledge to assemble in book form. This situation has changed dramatically for geriatric psychiatry in recent years. We hope that the knowledge gathered here from our distinguished international panel of authors bears this out.

John R. M. Copeland
Mohammed T. Abou-Saleh
Dan G. Blazer

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Part A

Historical Background

A Conceptual History in the Nineteenth Century

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The history of geriatric psychiatry can be written from two viewpoints. The “externalist” approach focuses on the social and political variables that have controlled attitudes towards abnormal behaviour in old age, and on the professionalization of those charged with the care of the mentally infirm elderly. The “internalist” approach—to be followed in this chapter—concentrates on the origin of the scientific language of psychogeriatrics. An adequate historical account should include information on theories of ageing, both physical and mental, brain sclerosis and the formation of a viable concept of mental illness. On the first rubric much research has been done¹⁻⁹; far less work exists on the other two. On psychogeriatric care before the nineteenth century^{10,11} there is very little: this may simply reflect a historical reality.

VIEWS ON AGEING BEFORE THE NINETEENTH CENTURY

Like most other aspects of human life, ageing has also been portrayed in terms of metaphors. Classical views, following the nature—nurture controversy, conceived of ageing as resulting from either internal instructions or from the buffeting of foreign factors^{4,8}.

The “wear and tear” view happened to be popular during the early nineteenth century, the period on which this chapter will concentrate. It was based, as it had always been, on the ageless observation that all natural objects, whether animate or not, are subject to the ravages of time. Surprisingly enough, the “wear and tear” view has not always generated an understanding attitude. In fact, across times and cultures great ambiguity has existed in regard to the treatment of old folk. Fortunately, a realistic acceptance seems to have predominated although there is plenty of evidence of hostility. The Hebrew tradition, and indeed its Christian offshoot, encouraged much reverence towards the wisdom and value of old age. But even in societies which have made great play of this view, veneration has been reserved for those in positions of power or influence¹². Little is known about attitudes towards elderly women or old men in humbler stations¹¹.

So, it can be concluded that, all in all, a view seems to have predominated that ageing was undesirable and that the identification of wear factors was important to devising ways of prolonging life^{5,13}.

A second ambiguity can be detected in these earlier writings. It concerns the extent to which the ageing process necessarily involves the human mind. Whilst it was a palpable fact that all human frames decayed, not everyone accepted that this had

necessarily to affect the soul or mind. Extant descriptions of the psychological changes brought about by old age suggest that people were aware that the mind also underwent a decline. However, theory and religion encouraged the view that the spirit could or did escape wear and tear, and that human beings grew ever more wise and useful, thanks to the accumulation of experience and knowledge. This belief must have been available in all those societies that felt the need to create adequate spaces for all manner of intellectual and/or sociopolitical gerontocracies². Some seem even to have separated chronological age and functional age in order to justify such concessions. From the point of view of the history of psychogeriatrics, it would be useful to know to what extent this belief was undermined by the occasional case of dementia amongst those elderly in positions of power¹. Historical evidence seems to show that these situations were neither more nor less perturbing than mental illness occurring at other periods of life. Indeed, fail-safe devices seem to have been available in these societies to cope with the upheavals created by such occurrences.

Men like Buffon, Darwin and Goethe reshaped ideas on ageing during the eighteenth century. Buffon¹⁴ wrote: “All changes and dies in Nature. As soon as it reaches its point of perfection it begins to decay. At first this is subtle and it takes years for one to realise that major changes have in fact taken place” (p. 106). Buffon put this down to an “ossification” process similar to that affecting trees: “this cause of death is common to animals and vegetables. Oaks die as their core becomes so hard that they can no longer feed. They trap humidity, and this eventually makes them rot away” (p. 111).

Erasmus Darwin’s views resulted from the application of yet another metaphor, namely, that ageing results from a breakdown of “communication” between man and his environment¹⁵. Darwin suggested that such breakdown followed a loss of irritability (a property of nerve fibres) and a decreased response to sensation:

“It seems our bodies by long habit cease to obey the stimulus of the aliment, which support us . . . three causes may conspire to render our nerves less excitable: 1. If a stimulus be greater than natural, it produces too great an exertion of the stimulated organ, and in consequence exhausts the spirit of animation; and the moving organ ceases to act, even though the stimulus is continued. 2. If excitations weaker than natural be applied, so as not to excite the organ into action, they may be gradually increased, without exciting the organ into action, which will thus acquire a habit of disobedience to the stimulus. 3. When irritative motions continue to be produced in consequence of stimulus, but are not succeeded by sensation . . .” (p. 365).

VIEWS ON AGEING DURING THE NINETEENTH CENTURY

In 1807 Sir John Sinclair¹⁶ published a major compendium on ageing and longevity which included references to most pre-nineteenth century sources. It was, in a way, the last grand glance to the past. Soon afterwards work started by those who, like Léon Rostan (1791–1866), based their claims on empirical findings. Rostan, one of the most original members of the Paris school, published in 1819 his *Recherches sur le Ramollissement du Cerveau*¹⁷, where the view commenced that vascular disorders might be as important as parenchymal ones in brain ageing. Even more important was his uncompromising anti-vitalistic position enshrined in the claim that all diseases were related to pathological changes in specific organs^{18,19}.

During the 1850s Reveillé-Parise³ saw his task as writing on “the history of ageing, that is, mapping the imprint of time on the human body, whether on its organs or on its spiritual essence” (p. v). In regard to ageing itself he wrote: “the cause of ageing is a gradual increase in the work of decomposition . . . but how does it happen? What are the laws that control the degradation that affects the organization and mind of man?” (p. 13). Reveillé-Parise dismissed the toxic view defended by the Italian writer Michel Lévy²⁰ according to which there was a gradual accumulation of calcium phosphates that led to petrification, to an “anticipation of the grave”. This view, he stated, had no empirical foundation and was based on a generalization from localized findings. Reveillé-Parise supported the view that ageing results from a negative balance between composition and elimination which equally affected the cardiovascular, respiratory and reproductive organs.

Finally, the views should be mentioned of J. M. Charcot, who in 1868 offered a series of 24 lectures on the diseases affecting the elderly²¹. Charcot dedicated Lecture 1 to the “general characters of senile pathology”; he started by saying that all books on geriatrics up to his time had “a particularly literary or philosophical turn [and had been] more or less ingenious paraphrases of the famous treatise *De Senectute*” (p. 25). He praised Rostan for his views on asthma and brain softening in the elderly, and predictably also mentioned Cruveilhier, Hourman and Dechambre, Durand-Fardel and Prus. He criticized Canstatt and other German physicians because in their work, “imagination holds an immense place at the expense of impartial and positive observation” (p. 26). Charcot’s own contribution was based on the general principle that “changes of texture impressed on the organism by old age sometimes become so marked, that the physiological and pathological states seem to merge into one another by insensible transitions, *and cannot be clearly distinguished*” (p. 27).

THE DEVELOPMENT OF THE NOTION OF BRAIN SCLEROSIS

When in 1833 Lobstein²² described the basic pathology of arteriosclerosis, he did not imagine that it would, during the second half of the century, become the mechanism of “senility” *par excellence*^{14,23,24}. Motor and sensory deficits, vertigo, delusions, hallucinations and volitional, cognitive and affective disorder were all attributed to the effect of arteriosclerosis^{25,26}. They related to the brain via a two-stage speculative pathophysiology: parenchymal and/or vascular disorders could affect the brain, and the distribution of the lesions could be diffused or focal. Vascular changes included acute ischaemia (on which clinical observation was adequate)^{27,28} and chronic ischaemia, invented as a separate syndrome by extrapolating from the symptoms and signs observed during the acute states²⁴. The role of arteriosclerosis as a causal and prognostic factor in relation to

the involuntional psychoses was challenged early in the twentieth century²⁹ but this paper remained unnoticed. Hence, some of the old notions, such as that of “arteriosclerotic dementia”, remained active well into the 1960s³⁰.

Alienists during the same period, however, were already able to distinguish between states where a putative chronic and diffuse reduction in blood supply had taken place from focalized damage, i.e. what they called “multifocal arteriosclerotic dementia” and was equivalent to what is currently called multi-infarct dementia^{24,31,32}.

NINETEENTH CENTURY VIEWS ON MENTAL DECAY IN THE ELDERLY

It is against this background that the history of the language and concepts dedicated to understanding mental disorders in the elderly must be understood. In addition to these neurobiological frameworks, a psychological theory that explained the manner of the decline was required. Such a psychopathology was provided by the heuristic combination of associationism, faculty psychology³³, and statistics³⁴ that characterized the early and middle part of the nineteenth century.

Yet another perspective, originating in clinical observation, was added during the 1830s. It led to the realization that, in addition to the well known forms of mental disorder, the elderly might exhibit specific forms of deterioration, and that these could be related to recognizable brain changes. There is only space in this chapter to deal with two examples: one typifying a “specific” disorder of old age, namely the history of chronic cognitive failure or dementia; the other illustrating the effect of a general mental disorder (melancholia) on the elderly.

THE FORMATION OF THE CONCEPT OF SENILE DEMENTIA

The history of the word and concept of dementia before the nineteenth century has been touched upon elsewhere³⁵. Suffice it to say here that, at the beginning of the last century, “dementia” had a “legal” and a “medical” meaning and referred to most acquired states of intellectual dysfunction that resulted in serious psychosocial incompetence. Neither age of acquisition nor reversibility was part of its definition. These two dimensions were only incorporated during the nineteenth century and completely changed the semantic territory of the dementia concept.

Anecdotal observation of cases of senile dementia abound both in the fictional literature and in historical documents³⁶, but the concept of “senile dementia”, as it is currently understood, only took shape during the latter part of the nineteenth century. Indeed, it could not have been otherwise, as the neurobiological and clinical language that made it possible only became available during this period^{37,38}. But even after the nosological status of senile dementia had become clearer, there were many who, like Rauzier³⁹, felt able to state: “it may appear either as a primary state or follow most of the mental disorders affecting the elderly” (p. 615). Following Rogues de Fursac⁴⁰, Adrien Pic—the author of one of the most influential geriatric manuals during this period⁴¹—defined senile dementia as: “a state of intellectual decline, whether or not accompanied by delusions, that results from brain lesions associated with ageing” (pp. 364–365). It was against this background that the concept of Alzheimer’s disease, which became the prototype for all senile dementias, was created during the first decade of the twentieth century³⁷. Recent work has shown that its “discovery” was controlled by ideological forces well beyond what could be described as “scientific”^{37,42}. These

forces also introduced unwarranted clinical strictures, such as the exclusion of non-cognitive symptoms^{43,44} and false age boundaries, which took many years to disappear.

THE FORMATION OF THE CONCEPT OF INVOLUTIONAL MELANCHOLIA

The concept of “senile or involutional psychoses”, which featured so prominently in Kraepelin’s early classification, included: pre-senile delusional insanity, senile dementia, late catatonia and involutional melancholia^{45,46}. The reasons that led Kraepelin to separate this group were mostly theoretical, to wit, that they appeared during a period of life when “sclerotic” changes were beginning to occur; the same factor accounted for their bad prognosis⁴⁶.

The general history of melancholia and depression has been analysed elsewhere⁴⁷⁻⁴⁹. Suffice it to say here that by the 1860s depression was considered to be an independent syndrome resulting from a primary disorder of affect. This meant that hallucinations, delusions and cognitive impairment were secondary to the pathological feelings. This conviction was particularly strong towards the end of the century, when emotional mechanisms became popular in the explanation of most forms of mental disorder⁵⁰. By the end of the century the metaphor of depression as a form of “reduction” or “loss” had become firmly established. No better example can be found than the fact that up to 1893 (fourth edition) Kraepelin felt obliged to classify all forms of agitated depression as mania⁵¹.

KRAEPELIN AND INVOLUTIONAL MELANCHOLIA

Much of the current confusion on the meaning of involutional melancholia can be explained if attention is given to the circumstances of its historical development (for a full analysis of this process and list of references, see reference 52). The conventional story⁵³⁻⁵⁶ is that up to the seventh edition of his textbook Kraepelin considered involutional melancholia as a separate disease, and that when confronted by the evidence collected by Dreyfus⁵⁷, he decided to include it, in the eighth edition, under the general heading of manic depressive insanity. Indeed, this account was first offered by Kraepelin himself (see reference 57, p. 169).

The story is, however, more complex and it is unlikely that the findings of Dreyfus alone caused Kraepelin’s change of heart. For example, Thalbitzer⁵⁸ claimed that his own work had also been influential (p. 41). In the eighth edition Kraepelin abandoned not only involutional melancholia but the entire group of “senile psychoses”. A recent statistical analysis of Dreyfus’s old series has also shown that his conclusion that the natural history of involutional melancholia was no different from that of depression affecting younger subjects was wrong⁵¹.

CONCLUSIONS

This short chapter, providing a historical vignette on the origin of the language of old age psychiatry, suggests that it was born during the nineteenth century from three conceptual sources: theories of ageing, neurobiological hypotheses concerning brain sclerosis, and the realization that specific forms of mental disorder might affect the elderly. Two clinical illustrations were provided, one pertaining to the origins of the concept of senile dementia, and the other to the notion of involutional melancholia.

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Scope and Development in the Twentieth Century

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Interest in the health and well-being of the elderly has existed since antiquity; over the centuries some remarkable observations were made regarding the health, the mental changes, and the care of the elderly. Some explanations were offered for age changes that were reasonable and some were fanciful, limited by existing scientific knowledge¹. During the twentieth century, many biological and behavioral theories of aging have been advanced and tested, emphasizing that aging is a multidimensional phenomenon.

To understand the rapid emergence of the psychiatry of old age during the twentieth century, one must appreciate that around the year 1900, the age composition of the population began to change. Life expectancies for both males and females were being extended, particularly for females. This was paralleled by a rapid expansion in science and technology.

Although the term “gerontology” has been around for many years, the term “geriatrics” was not coined until 1914 by Nascher. In that year he published the book *Geriatrics. The Diseases of Old Age and Their Treatment*². A prolific writer, Nascher’s last publication in 1944 was *The Aging Mind*³. He observed that chronic brain syndrome was likely to be familial and he believed that it was an accelerated primary aging process that was influenced by heredity. Seven years prior to Nascher, Alzheimer published his landmark report, which described what is now known as Alzheimer’s disease and is sometimes referred to as “the disease of the century”⁴. In the 1960s, books on aging and psychiatry began to appear. Among the best known of those in the English language were *The Clinical Psychiatry of Late Life*⁵ by Felix Post and *Behavior and Adaptation in Late Life*⁶. In the ensuing years, many important publications have appeared that reflect rapid advances in scientific knowledge and clinical practice. In the 1940s, gerontological and geriatric societies were organized in many countries in Europe and in North America. Early in 1950 the International Association of Gerontology was founded in London, UK, with the support of the Nuffield Foundation. The first international congress of gerontology was held in Liège, Belgium, July 9–12, 1950⁷. The International Psychogeriatric Association was founded in 1980 and held its first meeting in Cairo in 1982⁸.

GERIATRIC MEDICINE AND GERIATRIC PSYCHIATRY

For the past several decades, psychiatry and geriatrics have been experiencing what is called by many an “identity crisis”⁹. This crisis in identity centers on the sphere of professional activity that is the proper task of psychiatrists and geriatricians and whether their activity produces a source of self-esteem to the physicians.

The geriatric psychiatrist is making a significant contribution to solving this so-called identity crisis. The geriatric psychiatrist must often function as a primary care physician and his/her skills include proficiency in psychiatry, geriatric medicine, neurology and the social sciences. Geriatric psychiatrists are very aware that many of the disorders they treat can be cured or prevented but the resulting suffering can be relieved and the disability reduced. The situation does encourage the clinician to make observations that contribute to a better understanding of the course of chronic illness and to look for hidden clues that can lead to investigations which may, in the future, bring improved convalescence or even eradication of chronic disease and disability.

TECHNOLOGY

Advances in technology have widened the scope of old age psychiatry. For example, biochemical changes in the brain are distorted by the dying process and postmortem events. Routine postmortem examinations have limited usefulness, although rapid autopsies have improved the situation. *In vivo* brain imaging has enhanced our ability to observe biochemical processes and alterations in the aging brain¹⁰.

PSYCHOPHARMACOLOGICAL APPROACH

The psychopharmacological approach is currently the dominant treatment for mental and emotional symptoms in elderly people. Psychotropic medications often reduce or eliminate symptoms, but do not alter the cause of the disorder. Consequently, very critically needed and exciting opportunities for research lie ahead for the geriatric psychiatrist.

Because of the possible multiple etiology of many psychiatric disorders of late life, it is highly likely that the geriatric psychiatrist will have to have a broad knowledge base and therapeutic skills including the use of medications, psychotherapeutic techniques, and procedures to reduce risks inherent in the socioeconomic status and environment.

PSYCHOTHERAPY

It is unfortunate that in 1905 Freud expressed the view that patients “near or above the fifties” were not suitable subjects for psychoanalysis¹¹. For many years, this undoubtedly affected therapists’ attitudes towards all psychotherapy for the elderly. Fortunately, a number of prominent psychoanalysts challenged

this Freudian view and reported psychotherapeutic success with older patients¹². Beginning in the mid-twentieth century, a number of psychotherapeutic techniques were described and clinically evaluated for their effectiveness. Some are particularly applicable to the elderly psychiatric patient and include both behavioral and cognitive forms of psychotherapy. In Europe and North America psychotherapeutic approaches have been successfully used for elderly outpatients with depression and/or hypochondriasis.

THE SUBSPECIALTY OF GERIATRIC PSYCHIATRY

Geriatric consultants have served in the National Health Service in the UK for many years. A geriatric consultant is not a primary care physician and sees patients by referral, usually from general practitioners¹³. The geriatric consultant is likely to be hospital-based. Psychogeriatric long-term care beds are increasing in number and the responsibility is usually assigned to a consultant psychiatrist specializing in the psychiatry of old age. In 1985, the Royal College of Physicians of London conducted their first examination of candidates for a diploma in geriatric medicine. Although the Board of Examiners of the College includes representatives from general practice and from psychogeriatrics, the future of old age psychiatry as an area of specialization in the UK remains uncertain. In 1989 the British Department of Health recognized psychogeriatrics as an official subspecialty.

In Canada, the Royal College of Physicians and Surgeons in 1981 conducted examinations for special competence in geriatric medicine. Although geriatric psychiatry is not recognized as a subspecialty, the College has encouraged the development of programs in geriatric psychiatry. Since 1988 in the USA, the American Board of Internal Medicine, in collaboration with the American Board of Family Practice, has offered by examination a certificate of added qualifications in geriatrics. The examination is administered to candidates from both Boards at the same time in the same testing centers and the criteria for qualification are identical for both Boards¹⁴.

Geriatric psychiatry was the first subspecialty area for which the American Board of Psychiatry and Neurology offered an examination for added qualifications. The first examination was given in April 1991 to 661 candidates and the second year to 578 candidates. Since the first examination 3435 certificates have been issued¹⁵. The next examination was scheduled for the year 2000 and the first re-certification in 2000 as well. All added qualifications in geriatric psychiatry will be time-limited to 10 years. In most European countries an examination is not required for qualification in a medical specialty. However, Sweden is moving to the examination as a requirement.

Recently there has been a rapid development of professional organizations concerned with geriatric psychiatry. Two of the most active are the American Association of Geriatric Psychiatry and the International Association of Geriatric Psychiatry. The American association has made available geriatric psychiatry self-assessment.

For many years, it was believed that many physicians are reluctant to become involved in geriatrics. Numerous explanations have been offered, including relatively low monetary compensation, the lack of satisfactory treatment outcomes and lack of personal satisfaction and scientific challenge¹⁶. This view has rapidly changed and considerable interest and satisfaction is evident among medical students as well as physicians in training and practice. A recent study of those who completed geriatric fellowships in geriatric medicine or psychiatry and have now been in practice for at least 3 years found that 93% were satisfied with their career choice, 80% felt that they had maintained professional status and prestige, 71% were satisfied with their incomes and 96% found personal gratification in taking care of elderly patients¹⁷.

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The Development in Britain

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Psychogeriatrics, the psychiatry of old age, was born as a service activity some 25 years ago; in 1989 it became an official specialty in the National Health Service (NHS). Until the 1960s interest in the mental disorders of old age had been largely confined to research, but today special psychiatric services for old people are widely established, and the care of mentally ill old people is recognized as a major issue by professional workers, governments and the lay public, not only in developed countries but also in the Third World. Britain has led this movement.

ORIGINS

At the end of the 1960s, about half-a-dozen psychiatrists were running psychogeriatric services. Now there are some 450 and most health districts have such a service¹. This movement has the backing of government and of the relevant professional and voluntary bodies. The origins of this service specialty are five-fold:

1. Pressure of the increase in the numbers of the aged, particularly of the very aged.
2. Growth in psychiatry's capacity to treat conditions previously regarded as hopeless².
3. The movement of psychiatry from mental hospitals into people's homes, and into the general hospital.
4. The effectiveness of geriatrics in British medicine.
5. The writings and teaching in the 1960s of a small group of figures such as Sir Martin Roth and David Kay in Newcastle, Felix Post in London, and the pathologist Nicholas Corsellis at Runwell, on epidemiology³, clinical features and prognosis⁴⁻⁶ and pathology of the mental disorders of old age⁷.

In 1966 a paper from Newcastle⁸ emphasized the intertwining of physical, mental and social factors in the psychosyndromes of old age, and called for general hospital facilities for collaborative assessment of these complex disorders. Other writers were concerned with the possibly damaging effects of "misplacement" of old people in units of the "wrong" specialty, and Duncan Macmillan, a great psychiatric innovator, with his colleagues set up an early assessment unit and took a special interest in the assessment and care of old people^{9,10}.

THE PSYCHOGERIATRIC MOVEMENT

A symposium at the end of the 1960s¹¹ stands as a statement of what had then been achieved, and of the directions in which people were looking. Half-a-dozen younger psychiatrists were meeting as a "coffee house group", sharing their experiences of setting up local

services. They established contacts with government and with national bodies and they wrote, lobbied and spoke at innumerable meetings. Above all, they influenced younger colleagues¹².

In 1973 they formed a Group within the Royal College of Psychiatrists and in 1978, as numbers grew, the Group achieved the status of a Specialist Section. The Section has provided guidance on development of services, norms for staff and facilities and advice on changing issues. Successive chairmen of the Section have reflected the development of the specialty through three "generations" of workers.

By 1985–1986 some 250 psychiatrists were running local psychiatric services specifically for the aged¹³, in a variety of different styles but with a common philosophy; there have been many reports describing such services (reviewed in ref. 14). Currently, it is likely that some 450 consultant psychiatrists are primarily engaged in this work.

From the late 1970s, international networks were established, both personally and through bodies such as the World Health Organization, the Geriatric Psychiatry Section of the World Psychiatric Association, and the International Psychogeriatric Association. For a decade a course on psychogeriatrics has been run in Nottingham for the British Council, with participants from over 30 countries; versions of this have been "exported" to Australia, Israel, Poland and Portugal. An apparatus of education is established, and the Royal College of Psychiatrists sets standards for training. There are full professors of psychogeriatrics at some seven UK universities and academic posts are established in most medical schools.

A TYPICAL SERVICE

Principles for providing such services, and accounts of resources needed, have been the subject of publications by individual workers, by the Royal College of Psychiatrists and by the government (reviewed in ref. 14). *The Rising Tide* from the NHS Health Advisory Service¹⁵ has also been influential.

Developments have generally been around a group of core workers and facilities, the latter often determined by what happened to be available in the locality. One, and preferably two, psychiatrists will have special responsibility for the aged, working with nursing (including community nursing), remedial, social services, psychology and housing department staff, and with family doctors and their teams, with geriatric medical services and with voluntary and private facilities. The service generally deals with all forms of mental illness in old age, of which the commonest is depression, and the most exacting is dementia; but old age psychiatry spans virtually the whole of psychiatry—few problems

abate with age, and many become more intricate, with the admixture of physical and social problems.

The main thrust of a service is to maintain function, independence and choice. Staff strive to bring services to people in their homes, but hospital admission must be available for those who need it, as must a consultative service for other departments. Most services see patients initially at home and much follow-up takes place there or in the day hospital. Many units run special services, such as "memory clinics" or relatives' support groups, or collaborate in running stroke, continence or "orthogeriatric" services.

Standard facilities comprise an admission unit, which should be in the district hospital, outpatient clinics and day hospitals. A close relationship should exist with geriatrics, and in a few cases (especially where geriatrics and psychogeriatrics are located apart from each other) there are joint "psychogeriatric assessment units", in which psychiatrists and geriatricians collaborate. In Nottingham's Department of Health Care of the Elderly, geriatricians, psychogeriatricians and related staff have worked together in one department.

Longer-stay units, also offering respite admission, nowadays are smaller and more "domestic", and often close to their local communities. Much long-stay care is now private. Long-stay care remains the "Achilles heel" of the care of the elderly; the government's reaction to the report of its Royal Commission on Long-stay Care is still awaited¹⁶, whilst a series of National Required Standards on all aspects of long-stay care are currently about to go out to consultation¹⁷.

Wattis' analysis gives some support to the claim that services for older people provided by specialist psychogeriatricians are stronger than those provided by general psychiatrists who spend only a minority of their clinical time in old age psychiatry⁸. Such specialist services have significantly more consultant and non-medical staff time per thousand elderly served and are more likely to have acute beds on a district general hospital site and long-stay beds within the catchment area served. Their consultants are more likely to be involved in educational activities and to report research interests.

Voluntary Organizations

There has been a great growth of public awareness of the problems of mentally ill older people and of those who look after them. Bodies such as the Alzheimer's Disease Societies (now active in many countries) form a generally vigorous alliance with the professions as pressure groups, sources of information or support, or as fund-raisers.

Research⁸

There is nowadays hardly a relevant university department that is not concerned with the mental disorders of old age; biological research on the dementias has moved fast, as has clinical research and research on services and on carers. National and international collaboration in clinical and epidemiological studies is growing. A major longitudinal study of cognitive change in old people under the auspices of the Medical Research Council has been completed. The advent of journals such as the *International Journal of Geriatric Psychiatry* and *International Psychogeriatrics* reflects the clearer identity of the field and the growth of research activity.

Education and Entry to the Specialty⁴

Demands for teaching come from the health professions and beyond (e.g. police, clergy, architects, designers). Teaching in old

Table 3.1. What does a psychogeriatric service do?

Assessment
Diagnosis
Hospital liaison
"Rehabilitation"
Continuing care
Long-term and intermittent care
Support for carers
Planning
Advocacy, liaison and fund-raising
Other services
Voluntary
Private
Non-health professions
Government
"The public"
The media
Advice (e.g. financial, legal, "ethical")
Education
Research

age psychiatry is now common in the training of most of the health professions, and post-qualification courses are increasingly available. Day symposia abound. To enter the specialty in the UK, doctors need first to complete the 3 year basic training in psychiatry before taking the MRCPsych diploma of the Royal College of Psychiatrists, which will usually include 6 months in a psychogeriatric unit. Following the examination, higher trainees, in psychogeriatrics will spend 2 years in a psychogeriatric service and 2 years in general psychiatry, usually maintaining links with the former through clinical work or research and obtaining special experience (e.g. in geriatric medicine, neurology or management). Such training will qualify the doctor for a post as a psychogeriatrician, i.e. a psychiatrist devoting all or the bulk of his/her time to the psychiatry of old age. Higher trainees in general psychiatry may opt to spend a year in psychogeriatrics, and would then be qualified to take a "special interest" post, such as may exist in small districts. Many "doctors with domestic commitments" have trained part-time in psychogeriatrics.

Training arrangements will need to be adjusted for doctors who, having opted to do a year in psychogeriatrics, decide that they want this to become their main activity and so will wish to complete the full 2 years. In a new specialty there need to be clear standards and carefully monitored training, along with flexibility. Table 3.1 summarizes the range and scope of the work.

INTERNATIONAL DEVELOPMENTS

Psychogeriatric services are now in being in most developed countries, and there is activity in this field in the Third World too²⁰. The International Psychogeriatric Association is a thriving body, as is the World Psychiatric Association's Geriatric Psychiatry Section, which has lately published a series of broadly-based Consensus Statements on the content of the specialty, on Organization and on Education²¹.

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4a

The Development in the USA, 1600–1900

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There is a dearth of information about the care of mentally ill older persons in the USA from the seventeenth through the nineteenth centuries. Despite an extensive literature on the history of the various stages of the mental health movement during the period, little is known about the treatment of the aged, members of ethnic and religious minority groups, or of women of any age who became ill between 1600 and 1900. Much reported here refers to a largely white, male, New England data set. Virtually no evidence comes from the south or from west of the Mississippi.

In colonial times the mentally ill, along with other classes of dependents, were treated as a local responsibility, primarily placed with their own or other families. Clergy and physicians alike subscribed to a belief in demoniacal possession that was widespread in seventeenth century America. Few colonial doctors investigated other possible causes of mental disorders; no “scientific” theory for dealing with the mentally ill was yet in vogue.

Legal cases involving persons suffering from mental illness were decided on an individual basis. In 1639, the Massachusetts General Court was empowered to determine settlement for wandering individuals. Six years later, as a result of the need for such rulings, three men were selected to form a committee to consider provisions of a law for “disposing of inmates and settling impotent aged persons and vagrants”¹. It is probable that a significant minority of the people affected by this law were mentally ill, but the extant records do not yield adequate documentation to verify the incidence or treatment of such cases.

The “violent” insane among public dependents were ordinarily treated as common criminals. The harmless mentally ill were treated almost the same as other paupers. Records have been found as early as 1676 of the levy of a small tax on a village in Massachusetts to help one man build and maintain a small block house for his son, who was “bereft of his natural senses”².

In an effort to deal with its indigent and insane citizens, Boston in the late 1600s established the first almshouse in New England. Following English custom (which was to continue in America well into the nineteenth century), indigents and petty offenders were herded indiscriminately into this poorhouse—the sick and well, the able-bodied and impotent, and law-breaking and law-abiding, young and aged, “worthy poor” and vagrants, sane and insane. In 1736, the “Poor-House, Work-House, and House of Corrections of New York City” was built in Manhattan. Elsewhere, poor-houses and houses of corrections also served as repositories for the mentally ill. The founding of the Pennsylvania Hospital in 1752 (with its special section for the insane in the basement) and in 1773 the opening of the Virginia Eastern Asylum (the first American institution exclusively for mental patients) at Williamsburg, Virginia, provided crucial steps in a more humane treatment

orientation for the mentally ill. No special treatment for the elderly on account of their age, however, was instituted.

Besides attempts to effect reforms to help the insane during the Revolutionary period (c. 1765–1820), a few Americans challenged prevailing beliefs. Dr Benjamin Rush, who joined the staff of physicians at Pennsylvania Hospital in 1783, became known as the Father of American Psychiatry, in part because he advanced the humane and intelligent treatment of the insane. In addition to studying the effects of the moon on mental illness, Rush proposed that ill women and men engage in meaningful work, a forerunner of occupational therapy. He wanted patients to write down all that troubled their minds. Rush believed in kind treatment: asylums were to hire intelligent men and women to attend the patients. All visitors who had a disturbing effect on patients were to be excluded.

Rush’s views on mental health in late life are worth mentioning, because they anticipated later views and because so few of his contemporaries took an interest in the subject. Older people, he felt, were naturally protected against certain maladies; madness, he observed, tended to attack mainly between the ages of 20 and 50. In his *Medical Inquiries and Other Observations on Diseases of the Mind*³, Rush claimed that “the moral faculties, when properly regulated and directed, never partake of the decay of the intellectual faculties in old age, even in persons of uncultivated minds”. Potential, not just decline, characterized even those past 80 years of age.

Ironically, the “enlightened” view that older persons did not suffer as greatly as younger people from the scourges of mental impairment justified, for many Americans, the view that it was acceptable to treat elderly persons who did need help as “invisible lunatics”. This perception was reinforced by two trends. First, in the wake of the Revolution, a rising tide of humanitarianism dominated reformist thought in the early 1800s. Officials removed restraints in mental hospitals in the expectation that more humane treatment would facilitate recovery. This “moral treatment” concept dominated psychiatric practice for over 20 years until the 1830s, when it was abandoned. Karl Menninger⁴ attributes its decline to the influx of immigrants crowding the hospitals and the emergence of a new scientific perspective. Second, physicians such as Dr Weir Mitchell and Dr Pliny Earle assembled a battery of statistics to convince the medical profession that mental illness was incurable^{5,6}. If so, lifetime follow-ups would inevitably disclose recurrences of mental illness in a patient. This prognosis struck a death blow at moral treatment, and ushered in a long era of therapeutic nihilism. “Psychiatrists tended more and more,” claims Gerald Grob, America’s foremost scholar of the history of mental illness, “to disregard the psychogenic aspects of mental illness and to emphasize its somatic etiology”⁷.

In this context, there seemed to be little sense in doing much for the old. The elderly were discouraged from care in the state psychiatric hospitals, because they were thought, by virtue of their age, to be untreatable. If there were low probability that insanity could be cured even if detected early and treated aggressively, the elderly should not be allowed to take up valuable space that could be used to treat the curable. Therefore, custodial care for the elderly insane was the best that could be provided by families, almshouses, or prisons^{8,9}.

The elderly, nevertheless, represented a significant proportion of the institutionalized population. A comprehensive survey of insanity among the general population of Massachusetts conducted in 1854, for instance, revealed that insane people in asylums aged 60 and above made up 9.8% of the inmates. Nearly one out of every five insane persons in Massachusetts was over 60 years old. Despite such data, Rosenkrantz and Vinovskis¹⁰ maintain that the pervasiveness of insanity among the elderly was not recognized by nineteenth century physicians; the elderly insane remained the least likely age group to be institutionalized.

A lack of systematic record-keeping concerning mental health admissions of the aged prior to 1900 makes it hazardous to generalize about trends. Even so, based on nineteenth century data from Massachusetts, the following propositions seem warranted. On the one hand, admission rates fluctuated from one decade to the next, which suggests that those who ran institutions had considerable power to determine who could enter. On the other hand, men outnumbered women in asylums, probably because families found older women more deserving of support and less disruptive or threatening in behavior. Alcohol abuse, it is worth noting, was more likely to be cited as a diagnosis of insanity for males⁸.

Because of the prevailing nihilism, the dependent elderly were generally sent to local almshouses rather than mental institutions prior to 1890¹¹. Superintendents may have discouraged the admission of elderly patients, considering them a threat to the therapeutic mission of the hospital. However, records showed that patients over 70 were not troublesome and were kindly tolerated. They did receive less specific medical therapy than younger inmates. Curiously, given the rampant nihilism, what care they received apparently proved efficacious: more than half of those in institutions in their seventh decade were discharged due to recovery or improvement. The rest, however, usually stayed until death.

Because a significant proportion of mental patients aged in place, it is not surprising that the percentage of the elderly in institutions was greater than that in the general population. In 1880, there were 140 public and private mental hospitals caring for nearly 41 000 patients; 9300 were kept in almshouses; the rest were cared for in their own homes. The Census of 1880 showed 91 997 insane persons out of a total American population of 50 000 000; 52% were female, 71% native born, and 93% white. Responsibility for the aged insane was usually divided between local almshouses and mental hospitals. Between 1880 and 1890, insane persons constituted nearly a quarter of the total almshouse population. Between 1851 and 1890 nearly 10% of California's institutionalized insane were 60 years or older; in Arizona in 1900 the figure was only 1.71%; and in Massachusetts in the 1880s it was 12.1%. Most of the aged insane were said to be suffering from some form of senility.

The increasing use of the term "senility" to characterize impairments in mental health in later years signals the emergence of new scientific views of senescence. William James in *Principles of Psychology*¹² and George Beard in *American Nervousness*¹³ contended that the majority of the elderly invariably experienced a decline in mental faculties and a decreased ability to learn new materials and/or to adapt to changing circumstance. Charles Brown-Séguard hypothesized that the decline resulted from "diminishing action of the spermatoc glands". W. A. N. Dorland¹⁴ believed that people's "creativity" declined after age 60. Metchnikoff's attribution of old age to a chronic disease process is indicative of the pessimistic thought regarding the well-being of the elderly in general¹⁵. In view of such thought, it is not surprising that the elderly mentally ill were regarded as quite hopeless and undeserving of efforts at rehabilitation. Indeed, there was so little interest in this population that "geriatrics" did not become a medical specialty until the twentieth century under the influence of Dr I. L. Nascher¹⁶.

As basic processes of aging became a focus of scientific inquiry for a growing number of researchers, others tried to understand the etiology of more late-life disorders. Alois Alzheimer established (in a 1907 report) a histologic picture of the disease that would later bear his name. But there was not much interest in old age-associated dementia until increases in adult life expectancy started to change the demographic make-up of industrial societies¹⁷.

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In the Beginning

The Late Felix Post

In 1943, after a year's early training as one of the war-time refugees of the Maudsley Hospital, Professor Aubrey Lewis passed me on to Professor D. K. Henderson and the Royal Edinburgh Hospital for Nervous and Mental Diseases, where I initially worked in the private department. During one of his rounds, Henderson said to me: "Post, do you see all these old people here? Why don't you write 'em up?" This I obediently did, and my article appeared in the *Journal of Mental Science*¹. The article started by demonstrating that the admission rate of patients over 60 to the Royal Edinburgh Hospital had risen between 1901 and 1941 more steeply than the proportion of this age group in the Scottish population. Interestingly, at this early date, I had found no difficulties in the differential diagnosis of my colleagues' and my own patients. There were 22 senile, arterio-sclerotic and presenile dementia patients, 20 manic-depressive patients, 25 patients suffering from involutional or senile melancholia and 51 patients with schizophrenia. Assuming that the functional psychoses were the concern of general psychiatry, the rest of the paper dealt with the dementias and with an attempt to link the type associated with delusions and hallucinations to earlier personality characteristics. I noted that a high proportion of dementia admissions had been precipitated by terminal confusional states, and that of 111 patients admitted over the preceding 4 years with organic psychoses, only 23 were still occupying beds. I made the false prediction that in the future the main burden of the hospital services would be represented by the chronicity and survival of melancholic and paranoid patients. I did not anticipate that electroconvulsive therapy (ECT) and antidepressive drugs, while producing lasting recoveries in only 25% of cases, would make at least temporary discharge from inpatient care possible in most cases.

Aubrey Lewis was more farsighted. He had published, with a psychiatric social worker³ a paper describing the psychiatric and social features of the patients in the Tooting Bec Hospital for Senile Dementia, London, UK, and in 1946 predicted, in the *Journal of Mental Science*³ that ageing and senility would become a major problem of psychiatry.

After army service, I consulted Lewis about possible positions and he recommended me for the post of assistant physician at the Maudsley Hospital. I flattered myself that in me Lewis had seen a future brilliant psychiatrist, but was soon to be disillusioned. Even before the Bethlem Royal and Maudsley Hospitals were united in 1948, Lewis had conceived the idea of using some of the Bethlem beds to establish a unit for patients over the age of 60. After a heated discussion with the Bethlem matron, Lewis obtained agreement for the admission of senile patients to a hospital which, like the Maudsley, had previously admitted only patients thought to be recoverable. Uncovering his batteries, he asked me to take

on the development of this Geriatric Unit. Once again, I obeyed (to say without enthusiasm would be an understatement) and, right up to my retirement, I continued also to run a unit and outpatient clinic for younger adults.

A report in the *Bethlem Maudsley Gazette*⁴ demonstrated that both the Bethlem staff and I had "caught fire". The article started with a tribute to Professor Aubrey Lewis and his almost revolutionary idea of including experience in geriatric psychiatry within postgraduate training. The article went on to describe how patients over 60 had gradually infiltrated the Bethlem wards to emerge as a unit for 26 women and 20 men. The two wards were staffed by the same number of senior and junior nurses as the other adult wards, with two trainee psychiatrists changing every 6 months to other departments. There was one psychiatric social worker (PSW), later usually assisted by a trainee. The occupational therapy department had collaborated with the nursing staff to devise and carry out a daily occupational programme as well as socializing activities. The PSW ran a weekly afternoon of handicrafts, tea and talk near the Maudsley, where throughout my tenure I conducted a weekly follow-up and supportive clinic. The first year during which the unit had been in full swing was 1952, and it was recorded that during that year there had been 3.00 admissions to each geriatric bed compared to 3.74 admissions to each general psychiatric place. Patients who had been dementing, but whose home care was no longer possible had been excluded from admission, though not rigidly, as well as patients with recurring illnesses that had been adequately treated at the Bethlem-Maudsley or other hospitals. Of 133 patients, nine died, only four had to be transferred to their regional mental hospitals, seven were resettled in homes for the elderly, while 113 could be returned to family care. One year after discharge, information was successfully obtained about 121 of 124 cases. Seven patients had died, including one suicide of a woman who had discharged herself. Thirty ex-patients had to be readmitted to our or other hospitals, thirty-five were still outside hospital but by no means symptom-free, but 45 patients would be classified as recovered. These relatively favourable results were due to 89 patients having suffered from affective illnesses: 24 had symptoms associated with brain damage, 10 were mainly paranoid and 10 were regarded as having psychoneurosis. In spite of 4-6 weeks of conservative management 52 patients had to be given ECT. I concluded the article by pointing to research needs and by opining that with 30-40% of patients admitted to British mental hospitals being over the age of 60, training in the special problems of this age group was essential for all entrants to general psychiatry.

The history of the beginning would be incomplete without a brief account of further developments. My little textbook (rightly out of print) and publications on the long-term outcome of

affective, paraphrenic and schizo-affective illnesses were largely my own work, but many of the junior psychiatrists made contributions and they and clinical psychologists, as well as social workers, instigated their own researches. Many later made a name for themselves, and some became leading psychogeriatricians. Among them were Tom Arie, the late L. K. Hems, David Jolley, Robin Jacoby, David Kay, Kenneth Shulman and, last but certainly not least, Raymond Levy. After the Bethlem-Maudsley had accepted a district commitment and the admission of involuntary (sectioned) patients, Raymond Levy and my successor, Klaus Bergmann (not a Bethlem trainee), managed to move the Geriatric Unit to the Maudsley, so much closer to the patients' family homes. Raymond Levy succeeded in establishing an Academic Department of Old Age Psychiatry, which has continued to conduct research into the dementias of late life, that most important subject, previously neglected on account of

admission restrictions before the hospital abandoned its ivory tower to accept a district commitment. With similar developments elsewhere, psychogeriatrics became a world movement, and Sir Aubrey Lewis would be pleased.

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Part B

Normal Ageing

- BI Theories of Ageing
- BII Brain Ageing
- BIII Psychology of Ageing
- BIV Sociology of Ageing

General Theories of Aging

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INTRODUCTION

There is no satisfactory composite theory of aging, but numerous theories have been advanced to explain how and why living organisms age and die^{1,2}. Given the multidimensionality of human beings, many theories of aging, some familiar and some overlapping, have been developed. Theories of aging are usually grouped by biological, psychological, or sociological sciences. A comprehensive review of theories of aging is beyond the scope of this chapter. However, the selected theories that are to be reviewed are particularly relevant to the psychiatry of old age.

Depending on the discipline, the phenomenon of aging takes on different definitions. Biologic aging is made up of a number of undesirable processes. There are multiple processes of aging that result in a decline in efficiency of the organism and end in its death. Aging, particularly in the psychosocial sciences, often includes a desirable process of maturation, that is, acquiring a desirable quality such as wisdom.

THE BIOLOGICAL THEORIES OF AGING

Biological theories of aging can be broken down into two broad categories: the developmental genetic theories (primary aging) and the stochastic theories (secondary aging processes)^{3,4}. Primary aging refers to those declines in function that are genetically controlled, while secondary (stochastic) aging consists of random changes resulting from acquired disease and trauma. If the hostile events related to secondary aging could be prevented, life would be extended but, because of primary aging, decline and death are inevitable.

Deliberate Biological Programming

Hayflick and Moorhead⁵ made an important contribution to our understanding of cellular aging. They demonstrated that human mitotic cells would divide a finite number of times and then the cell culture would die. Numerous studies demonstrated that the “normal” cells have a memory for the number of duplications, which is believed to be encoded in the genetic material. In contrast, cancer cells have abnormal chromosomes and have the capacity for dividing endlessly.

The capacity for a programmed cell death that exists in mitotic (dividing) cells is also present in postmitotic (non-dividing) cells. Cells incapable of dividing, such as neurons, do age and die. Certain types of brain cells, such as those located in the

hypothalamus, appear to be much more vulnerable than others. In aging, not only are neurons lost, but there are alterations of neuronal synapses and networks⁶.

Apoptosis is a “physiological normal process by which multicellular organisms get rid of injured, infected or developmentally unnecessary cells”^{7,8}. Apoptosis is similar, if not identical, to a programmed cell death. The term is derived from Greek and means the dropping of petals from flowers or leaves from trees. The term “necrosis” is also commonly used to describe another kind of cell death. Necrosis is a type of cell death that is the result of the failure of nutrients to reach the cell. Death by necrosis is quite different from death by apoptosis. In necrosis the cells blow up, spill their contents and start an immune response. In apoptosis, however, the cells seem to condense, but develop blebs that separate and form “apoptosis bodies”. Apoptosis bodies undergo rapid phagocytosis; thus, in apoptosis the cells die with little observable reaction, while when they die of necrosis they produce a widespread inflammatory response.

Genetics of Human Aging

In humans and in many other animal species, females outlive males and this difference can be attributed to genetic factors, as the male has a Y and an X chromosome and the female has two X chromosomes. The small Y chromosome does not appear to contain sufficient genetic material for the normal development or well-being of a human. Its primary function is to provide male characteristics⁹. Chromosomes contain the vast majority of genes, but a few exist within the mitochondria. Mitochondrial genes are important to aerobic respiration and age changes. All mitochondrial genes are inherited from the mother¹⁰.

The Aging Clock

Miller provides a working definition of aging as follows: “Aging is a process that converts fit adults into frail adults with a progressive increased risk of illness, injury and death”¹¹. He favors an explanation for aging the existence of a single aging clock. This aging clock is in turn linked to many of the observed biological clocks that are seen in both humans and animals. Miller holds that a useful way of determining biological age will eventually emerge rather than depending upon chronological age. He supports the view that the genetic control of lifespan should be considered relatively minor. He believes that it influences lifespan at no more than 15–35%. Living in a favorable

environment and having favorable living habits influence most of the lifespan.

Inherent in our functioning are the biological rhythms observed at the hormonal levels. Many of the biological rhythms are synchronized by control centers within the brainstem. The hypothalamus plays a particularly important role in the losses of homeostatic mechanisms in the body. The responsible nuclei (clusters of cells within the hypothalamus) change with aging. These cells decline not only in number but also in efficiency. Sleep changes in late life are clearly associated with alteration of control centers within the brainstem. Van Gool and Mirmiran¹² propose that our biological rhythms become desynchronized as we age.

The Free Radical Theory

A free radical is often considered a molecular fragment, as it has an unpaired electron. It is unstable and highly reactive. Free radicals are ubiquitous in living substances and are produced by normal metabolic processes as well as by external causes such as ionizing radiation, ozone and chemical toxins. Free radicals have been linked to DNA damage, the cross-linkage of collagen and the accumulation of age pigments^{13,14}.

The Accumulation of Waste

With the passage of time, certain pigments such as lipofuscin accumulate in neurons and other cells. While there is no direct evidence that these pigments may be harmful to these cells, there is an association with the wear and tear of aging. Interestingly, the accumulation of lipofuscin is limited to the cells that are capable of dividing¹⁵.

The Immune System and Aging

The autoimmune theory of aging was proposed by Burnett¹⁶, Walford¹⁷ and Comfort¹⁸. It was suggested that a small number of immunologically competent cells may mutate in such a fashion as to lose their tolerance to their host antigen and subsequently give rise to a clone of “renegade” cells¹⁹, producing antibodies that might result in death or damage in a large number of cells, including neurons. “Anti-brain antibodies” are believed to be related to neuronal injury in senile dementia of the Alzheimer type. Autoimmunity to vascular antigens has also been reported²⁰.

The immune system is a complex network, but it has been found that restoring certain components can improve immunity. Interleukin-2 (IL-2) declines with age and it appears that the administration of IL-2 may retard the human aging processes²¹.

Caloric Restriction and Aging

It is well established that physical inactivity and overeating contribute to obesity, which in turn increases morbidity and mortality²². Determining the pathophysiology of overeating is being given considerable attention. It appears that oxidative stress associated with excess caloric intake results in damage that impacts on the process of senescence and various diseases common in late life. In contrast, a calorie-restricted diet in laboratory animals results in an increase in average lifespan. However, it is unclear whether further caloric restrictions in non-obese humans will add to life expectancy.

Other Biological Theories of Aging

The disposable soma theory holds that nature’s demand for reproduction takes precedence over a demand for longevity²³. Nature gives a priority to those traits that are inherent in the organism that favor reproductive success—traits that enhance their fecundity. Hence, somatic cells are disposed of after achieving reproductive success. However, it appears that the selection of traits that are favorable for reproduction may indirectly influence lifespan because the increased reserve capacity to carry an animal to and through a longer reproductive life also adds to the animal’s capacity to live longer.

The brain–body weight theory is based on evidence that the heavier the brain when compared to the weight of the body, the more likely the organism is to be inclined to longevity²⁴. This relationship of brain and body is called “the index of cephalization”. Naturally there are deviations from this correlation and this in turn has made the theory of limited validity.

Biophysicists have proposed a theory of aging that aims to explain why larger living organisms tend to live longer than smaller organisms. One explanation is that “lifespan tends to lengthen and metabolism slows down in proportion to the quarter power of the animal’s body weight”, or what might be called a “scaling theory” of aging²⁵. The theory is sometimes linked to the system of distributing nutrients. The rate of a heartbeat is relevant to the distribution of nutrients. For example, the elephant lives much longer than the chicken and has a much slower heart rate. However, the capillaries of elephants are the same size as those found in the chicken.

PSYCHOLOGICAL THEORIES OF AGING

Psychologists have accumulated a wealth of information regarding mental stability and change in late life. As in biology, this information has not been integrated into a viable comprehensive theory. The main areas that have been studied by psychologists can be placed in three broad categories: cognition, personality and coping mechanisms.

Cognitive Psychology

The term “cognition” subsumes the range of human intellectual functioning²⁶. Cognition is to perceive, to remember, to reason, to make decisions, to solve problems and to integrate complex knowledge. Measures of various types of cognition are influenced by chronological age, environment, task characteristics and other influences. With advancing age, individual differences in cognitive functions seem to increase. A comprehensive coverage of the studies on intelligence and memory in old age is beyond the scope of this chapter and will be presented at greater length in Section BIII, while basic concepts in those areas will be introduced in this chapter. In general, adults with high intelligence and education will show minimum decline in their performances with increasing age, while a significant decline is observed in adults with lower intelligence and age. However, older adults in general tend to perform less well in new or novel situations²⁶.

Loss of memory is a common complaint of old age and has received considerable attention by psychologists. There are several theoretical models of memory functioning. Such theories attempt to define various stages of information processing. Often attempts are made to distinguish short-term from long-term memory. Other theorists talk about primary, secondary and tertiary memory.

Research on intellectual functioning has been under way for many years and has been productive. Intellectual performance

seems to be strongly influenced by physical health. However, patterns of stability and change across the life cycle vary according to the ability that is being measured²⁶. Perlmutter²⁷ crystallizes the issue of psychological change and stability by positing a “multiprocess phenomenon conception”, as opposed to Baltes’ “dual process phenomenon conception” of development followed by decline²⁸. Perlmutter sees decline as neither inevitable nor universal and says that some cognitive skills may improve or may be acquired as one ages. However, as one reaches the point of “terminal drop”²⁹, which is a curvilinear decline related to the distance of death rather than old age itself, there will be a decline in intellectual functioning.

Schaie’s 30 stage theory of adult cognitive development attempts to formulate four cognitive stages in sequence. The first stage in childhood and adolescence is “acquisitive”, which is followed by the “achievement” stage in young adulthood, then by the “responsible and executive” stage in the middle-aged individual, and finally the “reintegrative” stage in old age. The shift is translated essentially from “What should I know?” to “How should I use what I know?” to “Why should I know?”

Ribot³¹ advanced the “cognitive regression hypothesis” which hypothesized that the structures first formed are the last ones to degenerate in old age. This has not been proved to be a constant feature, depending on what components of cognition are being studied. Essentially, no adult age differences have been found in conservation, egocentrism and concept attainment³². However, when it comes to constructing classification, young children and the elderly tend to have a holistic perception, while older children and younger adults are more analytic³³⁻³⁵. The fact that older adults have an easier time learning dated items and retrieving dated items follows Ribot’s law. In regard to free recall, older adults as well as younger children use motoric encoding and real-life objects and do not score as well as young adults, who perform better on standard memory tasks³⁶.

Personality Theories

Thomae and Lehr³⁷ have proposed an antistage theory of aging, where personality, development and adjustment are affected by the historical events throughout the life cycle. This theory is in partial conflict with the eight-stage theory of Erikson³⁸, which is a stage theory of ego development through the life cycle, culminating with the stage of maturity, as the elderly person may find either ego integrity through satisfaction with his past life, or despair and disgust over past failures. Bortwinick³⁹ has noted increasing cautiousness with advancing age, with the degree of cautiousness being influenced by the type of problem and its timing. Okun *et al.*⁴⁰ point out that cautiousness is not strictly an age effect, but that differences can be attributed to cohort influence.

According to Neugarten and Gutman⁴¹, people maintain their personality characteristics in late life. When personality changes occur, they appear to be related to losses, particularly those involving health and social support systems. Some sex differences are noted; men are more affiliative and more nurturant, women are more individualistic and more aggressive as they become older.

Costa and McCrae⁴², in their literature review on personality stability throughout the life cycle, also report that series of longitudinal studies show stability of personality traits in adulthood. The variables studied have included anxiety, introversion, conservatism, irritability or apathy. Costa and McCrae themselves have proposed five broad factors in personality traits: neuroticism, extroversion, openness to experience, agreeableness and conscientiousness. While there is stability of personality throughout the lifespan of an adult, there are generational

differences, secondary to cohorts. Personality changes may be very tightly woven with mastery and the ability to cope.

SOCIAL THEORIES OF AGING

Broadly, the sociological theories of aging can be broken down into those that examine the relationship of the older person to society and those that study the role and status of the elder. In their disengagement theory, Cumming and Henry⁴³ claimed that the withdrawal of the elderly from their previous societal roles with reduction in all types of interaction, essentially a shift of attention from the outer world to the inner world, was desirable and helped the elderly to maintain life satisfaction. With their exchange theory of aging, Homans⁴⁴ and Blau⁴⁵ also suggested that elderly people withdrew from social interaction. Ongoing social exchanges had become more costly in old age and therefore less rewarding.

In contrast to the disengagement theory, the activity theory^{46,47} proposed that activity contributes to health and life satisfaction. Undoubtedly, the selection of activities to be pursued by the elderly is limited by the decline that accompanies aging. However, remaining active is felt to be good for the elderly.

Neugarten and Gutman⁴¹ sought a compromise in the continuity theory, by noting that older adults tend to behave in a pattern established in their earlier life as they cope and make adaptive choices. At times the person may disengage and at other times remain active. Atchley⁴⁸ felt that the continuity theory was an illusive concept because aging produces changes that cannot be completely offset, so that there is no going back to a prior state.

Age and sex stratification provide different perspectives about aging by looking at different age and sex groups with different roles and expectations. As each group moves through time, it responds to changes in the environment⁴⁹. Riley⁵⁰ describes a cohort effect or “cohort flow”, where a group of people born at the same time in history are together and have certain common experiences and characteristics. The status of the aged is high in static societies and tends to decline with rapid social change⁵¹. This ties in with Cowgill and Holmes⁵² modernization theory, which suggests that the status of the aged in any society is inversely related to the level of industrialization within that society. With industrialization, the powers and prestige of the elderly are reduced. In a primitive society aging can be a liability, but older people who continue to perform useful and valued roles have a higher standing and are well treated⁵². These sociological theories have varying degrees of validity. In sociology, as well as in biology and psychology, there are no overarching theories that incorporate the theories described above.

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Structural Changes in the Aging Brain

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It has long been recognized that aging is associated with progressive changes in brain structure. The extent to which the brain is affected by aging is determined by a complex interplay of environmental and genetic factors. Little information is currently available on the influence of specific factors on brain aging; however, a significant amount of work has been done describing the aggregate of changes in the aging brain¹.

Methods applied to the study of brain aging have included anatomic studies and neuroimaging. Each method has specific strengths and weaknesses, leading at times to confusing and even contradictory results when comparing studies using different techniques. For instance, anatomic studies are subject to fixation artifacts and have rarely controlled for subjects' cranial size. On the other hand, magnetic resonance imaging (MRI) volumetry suffers from distortion effects due to imperfections in the magnetic field and from difficulty in precisely gauging the gray matter–white matter junction^{2–4}. Furthermore, no consensus exists regarding the definition of normal brain aging. Attempts at a definition have been confounded by various problems, such as difficulties in screening out individuals with subclinical neurodegenerative disease, and difficulties in conducting adequate longitudinal studies.

Despite these shortcomings, some information is available on the regional distribution of age-associated atrophy. A general principle of brain aging that has emerged is that phylogenetically newer structures, such as neocortex and neostriatum, tend to show more age-associated atrophy than phylogenetically older structures, such as the hippocampus and brainstem.

NEUROIMAGING STUDIES

Studies using various neuroimaging techniques have repeatedly demonstrated progressive age-related volume loss in widespread areas in the brain. The overall loss of tissue volume averages 0.2%/year, starting in mid-adulthood. Changes are manifested primarily by increases in ventricular volume and by cortical atrophy, as measured by increased sulcal markings^{9,10}.

These changes have significant clinical consequences. Increases in ventricular volume result in decreased cerebral compliance, rendering the aging brain less resilient to changes in intracranial pressure. Cortical atrophy, on the other hand, causes the brain parenchyma to pull away from the skull table, lengthening cortical bridging veins and increasing vulnerability to subdural hemorrhages.

In neuroimaging studies, gray matter volume loss has been found to be typically greater than white matter loss. Most affected is the pre-frontal cortex, with a rate of volume loss of about 0.5%/

year. Areas of moderate volume loss include the temporal, parietal and cerebellar cortex. The occipital cortex is relatively spared, while pons, tectum and hippocampus show minimal changes with normal aging⁹.

Various explanations for these differences in rate of atrophy have been offered, including regional differences in calcium homeostasis and expression of inflammatory mediators, such as 5-lipoxygenase. None of these hypotheses has been verified. Another important consideration is that distortions associated with image acquisition, such as movement artifact and inhomogeneities in the magnetic field, may result in insufficient precision to accurately measure volume changes in smaller structures, such as the pons.

The relative absence of hippocampal changes in normal aging stands in contrast to the dramatic decline in hippocampal volumes observed during the course of Alzheimer's disease, frontotemporal dementia and certain other disorders, such as mesial temporal sclerosis. Some researchers have attempted to exploit this difference as a means to assist in the diagnosis of Alzheimer's disease; however, hippocampal volumetry remains an experimental technique at present².

Aside from changes in volume, neuroimaging has revealed other interesting structural changes with aging. The most intensely studied of these changes is the increasing prevalence of leuko-araiosis, or periventricular white matter hyperintensities. The significance of leuko-araiosis is uncertain, but is thought to be related to tissue damage as a consequence of microvascular disease. Leuko-araiosis tends to be more prominent in individuals with hypertension or diabetes, although some degree of perivascular white matter changes are found in most individuals above age 65. To date, no strong relationship has been found between leuko-araiosis and cognitive decline^{3,4}.

Other noteworthy age-related changes include accumulation of iron in the striatum, deep cerebellar nuclei and motor cortex and the deposition of calcium in the pineal gland, choroid plexus and in the walls of the basilar and middle cerebral arteries. The clinical significance of these changes is also uncertain.

POST-MORTEM STUDIES

One important question that cannot be addressed by neuroimaging studies is the extent of actual cell loss occurring with brain atrophy. A sizable number of gross and microanatomical post mortem studies have been conducted to investigate this and other types of anatomical changes that occur in the brain with aging, although so far very few studies have attempted to compare neuroimaging with histologic data in the same individuals⁵.

Post-mortem volumetric studies have confirmed the findings from the neuroimaging literature that the frontal lobe shrinks proportionately more than the temporal, parietal and occipital lobes. In contrast to neuroimaging findings, however, is the observation that white matter volume loss appears to be greater than that found in gray matter. One possible explanation for this discrepancy is that aging seems to increase the water content of the gray matter–white matter junction, possibly distorting its apparent location on MRI imaging⁵.

Post-mortem ultrastructural studies have shed light on a variety of questions regarding changes at the microscopic and cellular level. Typical findings within normal aging brains include dilated perivascular spaces (Virchow–Robin spaces), mild demyelination, reactive gliosis and appearance of vacuoles and lipofuscin within neuronal and glial cell bodies. While many of these changes are attributed to age-associated “wear and tear”, the overall degree of white matter disorganization is mild when compared with that seen in arteriosclerotic brain disease^{5–7}.

An important finding confirmed in most ultrastructural studies is that most of the age-associated change in brain volume is attributable to decreases in the volumes of individual cells rather than to dropout of neurons and glia. Studies have repeatedly confirmed that the proportion of small cortical neurons increases with age. There is also some evidence for simplification of dendritic arborization with age⁵.

However, evidence also exists for at least a small degree of cell loss in various regions of the brain during normal aging, including parts of the substantia nigra, the locus coeruleus and the suprachiasmatic nucleus of the hypothalamus. Some cell loss has also been demonstrated in the cortex, specifically in layers 2 and 4⁶.

Other findings reported in ultrastructural studies or aging are that neurofibrillary tangles, but not amyloid plaques, appear in the CA1 cortex of the hippocampus and layer II of the entorhinal cortex. Aging also appears to affect the terminal first-order sympathetic axons, as manifested by swollen axons with neurofilamentous aggregates adjacent to the second-order sympathetic neuronal bodies².

SUMMARY

While there is no current consensus on the definition of normal brain aging, certain patterns have been tentatively identified. These include the observation that phylogenetically newer structures seem to atrophy at a faster rate than older structures such as the hippocampus, diencephalon and brainstem, and that most of the atrophy observed with normal aging can be attributed to cell shrinkage and dearborization, rather than to cell loss. The clinical consequences of these changes include decreased resilience

to changes in intracranial pressure and increased vulnerability to subdural hemorrhages. Areas of observed significant cell loss include the substantia nigra, locus coeruleus and suprachiasmatic nucleus of the hypothalamus.

Many important unanswered questions remain regarding normal aging, including:

1. What is the relationship between structural changes in brain tissue and age-related changes in other organ systems?
2. To what extent does apoptosis affect the brain in normal aging?
3. What are the environmental and genetic determinants of the rate at which the brain ages?
4. To what extent do gender differences affect cerebral aging?
5. To what extent does cell loss in the substantia nigra, locus coeruleus and suprachiasmatic nucleus impact on the changes in balance, gait, mood and sleep pattern that are observed with aging? Is cell loss within these areas truly “normal” or does it represent pathology with a high degree of penetration, such as might be seen with an environmental toxin or with a ubiquitous viral infection?

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Anatomy of the Aging Brain

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INTRODUCTION

Knowledge of the spectrum and extent of changes in brain morphology that occur with 'normal aging' is critical to any understanding of age-related illnesses, such as dementia. Prior to the recent introduction of advanced brain imaging techniques, post mortem studies served as the only source of information regarding the anatomy of the aging human brain. A common finding among them was a reduction of brain size with age, consisting initially (in the seventh decade) of atrophy of the gray matter (widening of sulci and thinning of gyri), followed by a decline in the volume of the white matter (in the eighth and ninth decades of life)¹. The cortical atrophy appeared to be most prominent in the frontal and parietal parasagittal areas and consisted of a decrease in the volume rather than in the number of individual neurons². Ventricular enlargement was also observed, but the degree of the dilatation was variable among the different studies^{1,3}.

There are several sources of error inherent to post mortem studies. Most of the studies included subjects with very different causes of death, which themselves can lead to a variety of changes in brain morphology; in particular, the cause of death may be quite different for young vs. elderly cohorts. Another disadvantage of the post mortem measurements is that the brain can be altered in unpredictable ways prior to tissue processing, as well as during the process of fixation^{4,5}.

Many of these problems have now been obviated by the development of imaging techniques, such as computed X-ray tomography (CT) and magnetic resonance imaging (MRI), which have provided an opportunity to examine the human brain *in vivo*¹⁰⁵. In this chapter, we critically review those brain imaging studies that have investigated age-related changes in brain anatomy. Any understanding of these data must begin, however, with a discussion of relevant methodological issues.

ISSUES OF METHODOLOGY

Study Design

Studies of the aging brain can be classified as either cross-sectional or longitudinal. Cross-sectional studies select subjects from all age groups and examine them during roughly the same time period. Subject recruitment is relatively easy and large amounts of data can be acquired and analyzed in a short period of time. Two main limitations are inherent to cross-sectional studies; the secular or generation effect and the survivor or cohort effect. The secular effect refers to differences among successive generations with

respect to variables that could influence brain anatomy. For example, differences between elderly and non-elderly cohorts with respect to nutrition or socioeconomic status could be associated with differences in brain morphology that have nothing to do with aging *per se*. The survivor effect refers to the biased selection of very healthy individuals, since those with underlying diseases usually die earlier and do not have the chance to be selected for study.

Longitudinal studies follow the same subjects with repeated examinations over time. Data acquisition is very labor-intensive and takes place over long periods of time. Longitudinal studies are limited by a period effect (changes over time in the methodology of the study) and an attrition effect (dropouts), in which the participants available at the end of the study may be quite different from the sample recruited initially.

Sample Selection

Age-related changes in brain morphology will obviously depend critically upon the criteria used to select the population under study. 'Normal' volunteers from the community may differ markedly from patients with 'clinically normal' brain CT or MRI scans with respect to variables that may influence brain anatomy. Medical and psychiatric illness, smoking, alcohol consumption, nutrition, education, environment, sex and height are just some of the variables that must be considered when comparing brain anatomy among individuals. Population heterogeneity is especially great among the elderly, and Rowe and Kahn⁶ have suggested a distinction between 'usual aging' (no clinically obvious brain disease) and 'successful aging' (minimal decline in neurobiologic function in comparison with young subjects). The increased variability of measures of brain anatomy seen in 'normal' elderly populations may be due in part to the relative mix of subjects with usual vs. successful aging.

Imaging Technology

Brain imaging techniques differ in their safety, sensitivity and anatomic resolution. One of the oldest methods for *in vivo* brain imaging is pneumoencephalography (PEG). PEG is unsuitable ethically for the study of the normal brain, due to its associated discomfort and high morbidity rate. The procedure itself may also directly affect the size of the cerebrospinal fluid (CSF) spaces⁷. These problems limit the application of this technique to patients with sufficient neurologic symptomatology to warrant such an

invasive procedure. As a result, the 'normality' of the sample in studies using PEG can be easily criticized.

With advances in computer technology, two new brain imaging methods have been recently introduced, computed X-ray tomography (CT) and magnetic resonance imaging (MRI). The major advantage common to both techniques is the opportunity to study both the CSF spaces and the brain parenchyma in great detail. Computed X-ray tomography is a relatively simple and inexpensive technique, with the advantage of short study acquisition times. The method is limited, however, by low spatial and density resolution, partial volume artifacts (the effect of the coexistence of multiple tissue and fluid components in the same volume unit) and bone-hardening artifacts (false elevation of brain CT numbers adjacent to the skull)^{5,8}. These technical problems have limited the ability of CT to quantitate brain changes with aging.

Magnetic resonance imaging takes advantage of the magnetic characteristics of tissue protons to study the brain. The main advantages of this technique are the excellent resolution of gray matter, white matter and CSF and the high sensitivity to disease processes, due to the fact that multiple factors (proton density, proton environment, etc.) contribute to the signal characteristics of each structure. The images are free of bone artifacts and the brain can be examined across all three major planes (i.e. axial, coronal and sagittal). Another advantage of MRI technology is that it does not utilize ionizing radiation. Disadvantages of MRI are the partial volume artifacts, the relatively longer study acquisition times and the fact that claustrophobic subjects cannot tolerate the examination due to the physical characteristics of the equipment^{9,10}.

Method of Image Analysis

Anatomical studies of the aging brain have used either qualitative or quantitative measures of brain morphology. In *qualitative studies*, raters employ various scales to examine the parameters of interest including, for example, the degree of cortical atrophy and ventricular enlargement. Qualitative ratings are relatively easy to use, are clinically relevant¹¹, do not depend on advanced instrumentation and sophisticated computations, and may display significant correlations with more quantitative methods¹². On the other hand, the accuracy of these ratings depends on the skill of the raters and, as such, it is difficult to compare the results of studies from different authors using different scales. In addition, the sensitivity and resolution of the scales are limited by the number of rating categories (usually three to five)¹³.

The *quantitative measurements* can be categorized as volumetric, planimetric (area measurement of regions of interest) or linear (distance measurement between points of interest) (Table 7.1). Volumetric methods are very accurate (according to validation studies on phantoms) and are very sensitive to changes

in brain size^{8,14,15}. The accuracy of the volume measurements is increased by obtaining relatively thin imaging slices (< 5 mm), by reducing the interscan gap (contiguous slices are preferred) and by scanning the entire extent of the structure of interest.

Planimetric studies have the advantage of being less labor-intensive than volumetric measures and, although they correlate reasonably well with volumetric measurements, at least for structures with regular shape^{12,15}, their overall accuracy and sensitivity is not as good. Thus, for a simple three-dimensional structure, a doubling of the volume will result in only a 1.5-fold increase in the area of a cross-section. For more complicated shapes with irregular configurations, the correlation of planimetric to volumetric measurements is much worse.

Linear measurements are the least accurate and least sensitive, primarily because they have a non-linear relationship to volume measurements. As a result, small but potentially important changes in volume may be underestimated by linear measurements, even when used in combination¹⁵. Finally, it is important to note that the size of various brain structures or regions might be expected to vary with the size of the subject's brain; e.g. subjects with larger brains might be expected to have larger ventricles. Any morphometric study of brain structures across individuals should therefore take into consideration intergroup differences in brain size¹⁵⁻¹⁷.

LITERATURE REVIEW

This section summarizes the effects of aging on CSF space size, size of brain parenchyma, ventricular size, and incidence of subcortical hyperintensities as reported in the literature.

Table 7.2 reviews the reported changes in the size of brain parenchyma with aging. The proportion of sulcal CSF volume to cranial volume (an indirect measure of cortical gray matter atrophy) increases from approximately 3% at the second decade of life to approximately 10% at the ninth decade. This age-related increase in sulcal CSF volume appears to accelerate after the sixth decade. Studies that have directly measured brain parenchyma have reported a 15-25% decrease in cortical gray matter volume, while the white matter, in most studies, appears to remain stable. However, studies of the corpus callosum consistently demonstrate age-related reductions in total callosal volume, particularly in the anterior regions. Finally, age-associated reductions in the areas of the pituitary and cerebellar vermis, as well as in the volume of the caudate nucleus, have been reported, though attempts at replicating those results have yielded some negative findings.

Some studies have attempted to estimate reductions in brain size indirectly by measuring changes in the volume of CSF spaces (Table 7.3). Results consistently indicate that the size of CSF spaces increases with age. More specifically, ventricular:brain ratio (VBR) may increase in a non-linear manner from 2% to

Table 7.1. Linear measurements—definitions

<i>Bifrontal ratio</i>	Distance between the tips of the frontal horns divided by the distance between the inner tables of the skull. Measured at the slice best showing the caudate ¹⁹
<i>Bicaudate ratio</i>	Minimal distance between the caudate indentation of frontal horn divided by the distance between the inner tables of the skull. Measured at the slice best showing the caudate ¹⁹
<i>Lateral ventricular ratio</i>	Distance between the lateral walls of the bodies of the lateral ventricles divided by the distance between the inner tables of the skull. Measured at the slice showing the maximum area of bodies of lateral ventricles ¹⁶
<i>Cortical sulci ratio</i>	Sum of widths of the four widest sulci divided by the transpineal inner table diameter. Measured in two supraventricular slices cutting through the centrum semiovale ¹⁹
<i>Interhemispheric fissure ratio</i>	Maximal width of interhemispheric fissure divided by the transpineal coronal inner table diameter ¹⁹
<i>Sylvian fissure ratio</i>	Average of maximal Sylvian fissure width divided by the transpineal inner table diameter ¹⁹
<i>Third (III) ventricle ratio</i>	The product of the sagittal and coronal diameters of the IIIrd ventricle, divided by the product of the transpineal and midsagittal inner table diameters; measured at the level of maximum IIIrd ventricle area ¹⁹

Table 7.2. Aging and changes in size of brain parenchyma

Study	Subjects	Imaging and measurement technique	Findings
Haug 1977 ¹⁸	170 Scans 0–75 years old. Subjects with 'normal neurological findings' complaining of headaches	CT, linear (maximum width of interhemispheric fissure)	Cortical atrophy increased with age; width of interhemispheric fissure increased approximately 5-fold (0.5–2.8 mm) from age 16–30 to age 61–75, in a continuous fashion
Jacoby <i>et al.</i> , 1980 ¹⁹	50 Healthy elderly volunteers, 62–88 years old, 10 M, 40 F. No history of significant psychiatric or neurologic illness. Handedness not specified	CT. Ratings (four-point scale) of cortical atrophy from films by single blinded rater; five regions rated (frontal, parietal, temporal, insular and occipital) and scores summed	Age correlated with total cortical atrophy score. Interactions with sex or laterality not reported. Adjusting for age, no relation between any CT measure and performance on the Hodkinson test of memory and orientation
Cala <i>et al.</i> , 1981 ²⁰	115 Volunteers, 15–40 years old, 62 M, 53 F. No history of migraine, head trauma, or excessive alcohol intake (no additional details provided). All but eight subjects right-handed	CT (n = two scanners). Ratings (five-point scale) of cortical atrophy (no additional details provided). Axial slices (13 mm thick)	Age apparently associated with increased frequency of mild (grade 2) atrophy of frontal lobes and cerebellar vermis, but no statistical analysis reported. Interactions with sex or laterality not reported
Zatz <i>et al.</i> , 1982 ¹²	123 Normal volunteers, 20–90 years old. Unequal sex distributions among different age groups, with more women in the old group. Excluded: history of neurological problems or major medical diseases	CT. Planimetry (sulcal fluid area in two supraventricular slices divided by cranial size). Volumetric (total fluid volume in temporo-Sylvian area divided by total cranial volume). Axial slices (n = 9) 10 mm thick, with 10 mm interscan gap; lowest slice at the level just above the petrous pyramids and orbital roofs	Increased cortical atrophy with age. Area of sulcal fluid in supraventricular levels was stable until the seventh decade then increased four-fold from age 60 to 80 +. Volume of fluid in temporo-Sylvian area was stable until age 60; then increased by 30% from age 60 to age 80 +
Gado <i>et al.</i> , 1983 ¹⁴	12 normal volunteers 64–81 years old. Excluded: subjects with dementia	CT. Sulcal volumetric ratio (ratio of sulcal to cranial volume). Axial slices (n = 7), including three sections above the roof of the lateral ventricle and four below it	Increased cortical atrophy with age; cortical sulcal volume ratio increased by 13.3% in 1 year follow-up
Gomori <i>et al.</i> , 1984 ²¹	148 Neurologically intact patients (subjects with minor neurological symptoms and patients with lung, breast, prostate or colon cancer but no CNS involvement. All subjects had normal neurological examinations), 28–84 years old	CT. Linear (cortical sulci ratio, frontal interhemispheric fissure ratio, Sylvian fissure ratio)	Cortical atrophy increased with age; age correlated with Sylvian fissure, interhemispheric fissure and cortical sulci ratios (r = 0.53, 0.4 and 0.2, respectively). Sylvian fissure ratio increased 1.6-fold between ages 28–49 and 80–84 continuously; interhemispheric fissure ratio increased five-fold across the same age groups continuously
Laffey <i>et al.</i> , 1984 ¹¹	212 Normal volunteers living independently, 65 years old and older. Sex distribution same in all groups (52% M, 48% F). Excluded: alcoholism and history or findings of neurological illness	CT. Qualitative rating (five-point scale) of cortical atrophy by two experienced radiologists. Axial slice at the mid-ventricular level	Trend (not significant) for increased cortical atrophy with age; mean rating of 0.83 at age 65–69 to 1.4 at age 80–89. Males showed higher scores for cortical atrophy in all age groups
Schwartz <i>et al.</i> , 1985 ⁵	30 Healthy male volunteers 21–81 years old. No history of major medical, neurologic or psychiatric illness. Handedness not specified	CT. Volume measurements derived from computer-assisted segmentation technique (ASI-II program). Axial slices (n = 7) starting from the plane of the inferior orbitomeatal line (10 mm thick, 7 mm interscan gap)	Adjusting for intracranial volume (IV), age negatively correlated with volume of gray matter and with volume of gray plus white matter, but not with white matter volume. Subjects more than 60 years old (n = 11) had smaller volumes of thalamus, lenticular nuclei and total gray matter than younger subjects (n = 19). Effects similar for both hemispheres
Pfefferbaum <i>et al.</i> , 1986 ¹³	57 Normal volunteers 20–84 years old, 27 M, 30 F	CT. Sulcal volumetric ratio (ratio of total sulcal volume to total cranial volume). Axial slices (n = 12) with 8 mm interscan gap, starting from the level of the superior roof of the orbit and proceeding upwards	Increased cortical atrophy with age; volume ratio increased seven-fold from the third to the eighth decade. The increase was faster after age 60
Stafford <i>et al.</i> , 1988 ²²	79 Normal male volunteers, 31–87 years old. Excluded: alcoholism, psychiatric illness, learning disability, severe head trauma, epilepsy, hypertension, chronic lung disease, renal disease, coronary artery disease, cancer	CT. Planimetry: ratio of cortical sulcal area to cranial area. Axial slice at supraventricular levels	Cortical atrophy increased with age; cortical sulci area ratio increased 1.8-fold between ages 30–39 and 70 + (0.012–0.022), faster after the sixth decade
Golomb <i>et al.</i> , 1993 ²³	154 Healthy volunteers with MMSE > 27, 55–88 years old (70 + 8 years), 73 M, 81 F. No evidence of active medical, neurologic, or psychiatric illness. Handedness not specified	CT (n = 51); MR imaging (n = 81); both CT and MR imaging (n = 22). Blinded ratings (4-point scale) of hippocampal atrophy as defined by dilatation of transverse choroidal fissure on films, by raters (n = ?) with established reliabilities	Subjects with hippocampal atrophy (rating of 2 or greater in either hemisphere; n = 50) significantly older than those without atrophy. More males (41%) than females (25%) with hippocampal atrophy

Table 7.2 continued

Study	Subjects	Imaging and measurement technique	Findings
Meyer <i>et al.</i> , 1994 ²⁴	81 Healthy volunteers, 27–90 years old, 44 M, 37 F. No major neurologic or psychiatric illness	CT ($n=2$ scanners). Blinded measure of tissue density (densitometry) and regional brain volume (trace methodology) from axial slices (8 mm thick)	Age associated with decreased tissue density in cortical gray matter (frontal, temporal, parietal, and occipital) and in white matter (frontal only), but not in subcortical gray matter (caudate, putamen, or thalamus). Age associated with decreased ratios of cortical gray matter volume to IV and subcortical gray matter volume to IV, but not with white matter volume to IV. Interactions with sex or laterality not reported
Elwan <i>et al.</i> , 1996b ²⁵	88 Healthy 'lower middle class' volunteers, 40–76 years old (54.8 + 9.6 years), 57 M, 31 F. No major medical, neurologic, or psychiatric illness. All right-handed	CT. Multiple linear measurements (no additional details provided)	No correlation between age and maximal bifrontal distance, bifrontal index, maximal bicaudate distance, maximal septum–caudate distance, or cella media index. Interactions with sex or laterality not reported
Yoshii <i>et al.</i> , 1986 ²⁶	33 Normal volunteers, 24–82 years old	Magnetic resonance (MR) imaging. Planimetry	No significant correlation between corpus callosum area and age. Trend for decline in the area of anterior half of corpus callosum with age ($r=0.31$, $p=0.8$)
Simon <i>et al.</i> , 1987 ²⁷	48 Subjects, including normal volunteers and patients with normal MR imaging studies, 11 months to 64 years old, 25 M, 23 F. Excluded: periventricular high signal areas, anomalous development of the corpus callosum, pre-scan diagnosis of multiple sclerosis	MR imaging (0.15 tesla). Planimetry; midsagittal slice, 5 mm thick	Corpus callosum area did not change after the 18th year of age
Uematsu <i>et al.</i> , 1988 ²⁸	17 normal male volunteers, mean age 31.5 years old, SD 5.5 years. Physically healthy	MR imaging (0.5 tesla). Planimetry; midsagittal slice 10 mm thick	No significant correlation between corpus callosum area and age
Condon <i>et al.</i> , 1988 ²⁹	40 Volunteers, 20–60 years old, 20 M, 20 F. No additional details provided	MR imaging (0.15 tesla). Volume measurement (two raters) derived from computer-assisted pixel segmentation of contiguous sagittal slices (variable slice thickness and number)	For males but not females, age negatively correlated with ratio of total brain volume to IV. Interactions with laterality not reported
Yoshii <i>et al.</i> , 1988 ³⁰	58 Volunteers, 21–81 years old, 29 M, 29 F. Neurologic and psychiatric histories not reported. Handedness not specified	MR imaging (1.0 tesla). Mathematically derived estimate of brain volume from inversion recovery films, based on planimetric area measurement made on single slice (10 mm thick) at level of foramen of Monro. Blinded global ratings of cortical atrophy from films (axial slices, [$n=?$], 10 mm thick, 3 mm interscan gap). Number of raters and rater reliabilities not specified	No correlation between age and brain volume. Age significantly correlated with ratings of cortical atrophy for both males and females
Hayakawa <i>et al.</i> , 1989 ³¹	143 Patients and seven normal volunteers, 0–60 years old. All with normal MR imaging scans	MR imaging (0.35 tesla). Planimetry; midsagittal slice 10 mm thick	Decrease in size of brain structures with age. Pituitary area decreased progressively after 20 years of age. Trend for decrease in cerebellar vermis area after the fifth decade; pontine and corpus callosum area remained stable during the adult life
Hauser <i>et al.</i> , 1989 ³²	25 normal volunteers, mean age 37 years old, SD 8.6 years, 14 M, 11 F. Excluded: history of medical or psychiatric illness	MR imaging (0.5 tesla). Planimetry; midsagittal section	Corpus callosum area did not correlate with age
Jernigan <i>et al.</i> , 1990 ³³	58 Healthy volunteers, 8–79 years old, 35 M, 23 F. No history of neurologic, psychiatric, or medical illness (diabetes mellitus, heart disease). Handedness not specified	MR imaging (1.5 tesla). Volume estimates (one of two raters) derived from computer-assisted pixel classification of multiple spin-echo axial images (5 mm thick, 2.5 mm interscan gap)	Age negatively correlated with ratios of cerebral volume to IV and of gray matter volume to IV. Among gray matter structures, age negatively correlated with ratios of cortical gray matter volume to IV, caudate volume to IV, and dicephalon volume to IV; but not with thalamus volume to IV or anterior cingulate volume to IV. No correlation between age and ratio of white matter volume to IV. Interactions with sex or laterality not reported

continued

Table 7.2 continued

Study	Subjects	Imaging and measurement technique	Findings
Krishnan <i>et al.</i> , 1990 ³⁴	39 Healthy volunteers, 24–79 years old, 17 M, 22 F. No evidence of major medical, neurologic, or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Stereological measurement (one of two raters) of axial slices (variable number, 5 mm thick, 2.5 mm interscan gap) from intermediate and T ₂ -weighted films	Age negatively correlated with total caudate volume (males = females). Caudate volume was less in subjects older than 50 years ($n = 22$). No adjustments for cranial size
Doraiswamy <i>et al.</i> , 1991 ³⁵	36 Healthy volunteers (overlap with subjects Krishnan <i>et al.</i> , 1990), 26–79 years old, 16 M, 20 F. No evidence of major medical, neurologic, or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Area measurement of T ₁ -weighted midsagittal image using computer-assisted trace methodology. Rater reliabilities not reported	Age negatively correlated with corpus callosum area in males but not in females
Escalona <i>et al.</i> , 1991 ³⁶	37 Healthy volunteers (overlap with subjects Krishnan <i>et al.</i> , 1990), 24–79 years old, 16 M, 21 F. No evidence of major medical, neurologic, or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Stereological measurement (one of two raters) of axial slices (variable number, 5 mm thick, 2.5 mm interscan gap) from intermediate and T ₂ -weighted films. Good rater reliabilities	No association between age and volume of cerebellar hemispheres
Gur <i>et al.</i> , 1991 ³⁷	69 Healthy volunteers, 18–80 years old, 34 M, 35 F. No neurologic or psychiatric illness; 66 right-handed; three left-handed	MR imaging (1.5 tesla). Volume measurements (any two of four raters) derived from segmentation technique based on two-feature pixel classification of multiple spin-echo axial images (5 mm thick, contiguous)	Older (555 years) subjects ($n = 26$) had smaller whole brain volumes than younger subjects (males = females)
McDonald <i>et al.</i> , 1991 ³⁸	36 Healthy volunteers (subjects also included in Krishnan <i>et al.</i> , 1990), 24–79 years old, 13 M, 23 F. No evidence of major medical, neurologic or psychiatric illness	MR imaging (1.5 tesla). Same as Krishnan <i>et al.</i> , 1990 (above)	Age negatively correlated with total putamen volume (males = females; left = right), but no adjustments for cranial size
Shah <i>et al.</i> , 1991 ³⁹	36 Healthy volunteers (overlap with subjects in Krishnan <i>et al.</i> , 1990), 26–79 years old, 16 M, 20 F. No evidence of major medical, neurologic, or psychiatric illness	MR imaging (1.5 tesla). Computer-assisted measurements from T ₁ -weighted midsagittal films by single rater with established intra-rater reliabilities	Increasing age associated with decreasing midbrain area (males > females?). No age effects on areas of pons, medulla, anterior cerebellar vermis or 4th ventricle
Tanna <i>et al.</i> , 1991 ⁴⁰	16 Healthy volunteers, 52–86 years old, 5 M, 11 F. No evidence of major medical, neurologic, or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Volume measurements (one of two raters with established reliabilities) derived from segmentation techniques based on two-feature pixel classification of multiple spin-echo axial images (5 mm thick, 2.5 mm interscan gap)	Age negatively correlated with ratio of total brain volume to total CSF plus total brain volume. Interactions with sex or laterality not reported
Coffey <i>et al.</i> , 1992 ⁴¹	76 Healthy volunteers, 36–91 years old, 25 M, 51 F. No lifetime history of neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla). Volume measurements (one of three blinded raters with established reliabilities) using computer-assisted trace methodology of T ₁ -weighted coronal images ($n = 30–35$, 5 mm thick, contiguous). Blinded clinical ratings (five-point scale) of 'cortical atrophy' (average score of two raters)	Age associated with decreased total volumes of the cerebral hemispheres (0.23%/year), the frontal lobes (0.55%/year), the temporal lobes (0.28%/year), and the amygdala–hippocampal complex (0.30% per year); all effects similar for males and females, and for both hemispheres. Increasing age associated with increasing odds (8.9%/year) of 'cortical atrophy', from 0.08 at age 40 to 2.82 at age 80
Doraiswamy <i>et al.</i> , 1992 ⁴²	75 Healthy volunteers (overlap with subjects in Krishnan <i>et al.</i> , 1990), 21–82 years old (52.5 + 18 years), 34 M, 41 F. No neurologic or psychiatric illness	MR imaging (1.5 tesla). Blinded stereological measurements of volume and linear measurements of size of midbrain on T ₂ -weighted axial films (no additional details provided)	Age negatively correlated with midbrain volume and anteroposterior diameter, but not with red nucleus size. Effects similar for both males and females
Jack <i>et al.</i> , 1992 ⁴³	22 Healthy elderly volunteers, 76.3 + 11.3 years old, 10 M, 12 F. No major medical or neurologic illness; no depression. Handedness not specified	MR imaging (1.5 tesla). Volume estimates (single rater) derived from computer-assisted pixel classification of T ₁ -weighted coronal images (4 mm thick, contiguous). Intra-rater reliabilities not reported	Age associated with decreased ratio of hippocampal volume to IV and of anterior temporal lobe volume to IV. Interactions with sex or laterality not reported
Lim <i>et al.</i> , 1992 ⁴⁴	14 Healthy male volunteers, eight young (21–25 years old) and six elderly (68–76 years old). No evidence of significant medical or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Blinded volume measurements derived from semi-automated pixel segmentation of intermediate and T ₂ -weighted axial images ($n = 8$, 5 mm thick, 2.5 mm interscan gap)	Compared to younger males, older males had lower ratio of gray matter volume to IV (49.7% vs. 38.7%). No group difference in ratio of white matter volume to IV (47.2% vs. 41.2%). Interactions with laterality not reported

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Table 7.2 continued

Study	Subjects	Imaging and measurement technique	Findings
Murphy <i>et al.</i> , 1992 ⁴⁵	27 Healthy male volunteers, 19–92 years old. No major medical, neurologic or psychiatric illness. Handedness not specified	MR imaging (0.5 tesla). Blinded volume measurements using computer-assisted trace methodology of proton density axial images ($n = 36$, 7 mm thick, contiguous). Manual tracing of subcortical from enhanced images. Rater reliabilities were established, but number of raters not reported	Older males (> 60 , $n = 17$) had smaller ratios of total, left, and right hemisphere volume to IV than younger males. Older males had smaller ratios of total caudate volume to IV and of total lenticular nuclei volume to IV than younger males; no difference in ratio of total thalamus volume to IV. Reductions in caudate and lenticular volumes also found when the volumes were normalized to total brain volume, suggesting a differential effect of aging on these structures. Older males exhibited a R > L asymmetry in lenticular nuclei; the reverse was true in younger males.
Doraiswamy <i>et al.</i> , 1993 ⁴⁶	Same as Doraiswamy <i>et al.</i> , 1992	MR imaging (1.5 tesla). Blinded linear measurements of interuncal distance on T ₁ -weighted axial image (no additional details provided)	Age associated with larger interuncal distance (NB: this measure was not correlated with amygdala volume in a follow-up study [Early <i>et al.</i> , 1993]). Interactions with sex or laterality not reported
Raz <i>et al.</i> , 1993a,b,c ^{47,48,49}	29 Healthy volunteers, 18–78 years old (43.8 + 21.5 years), 17 M, 12 F. No history of medical, neurologic, or psychiatric illness. All right-handed	MR imaging (0.3 tesla). Volume measurements using computer-assisted trace methodology of digitized images from the films, by two blinded raters with high reliabilities. T ₁ -weighted axial slices ($n = 9$, 4.2 mm thick, 6.0 mm interscan gap). T ₂ -weighted coronal slices ($n = 17–21$, 6.6 mm thick, 8.6 mm interscan gap)	After controlling for head size, age associated with decreased volumes of caudate and visual cortex (females > males). No association between age and volumes of dorsolateral prefrontal cortex, anterior cingulate gyrus, prefrontal white matter, hippocampal formation, postcentral gyrus, inferior parietal lobule or parietal white matter
Christiansen <i>et al.</i> , 1994 ⁵⁰	142 healthy volunteers, 21–80 years old, 78 M, 64 F. No major medical or neurologic illness	MR imaging (1.5 tesla). Area and volume measurements using computer-assisted trace methodology of T ₂ -weighted axial slices ($n = 15$, 4 mm thick, 4 mm interscan gap). Number of raters, their 'blindness' and their reliabilities not specified	Age associated with decreased volume of cerebral hemispheres. Interactions with sex or laterality not reported
Cowell <i>et al.</i> , 1994 ⁵¹	130 healthy volunteers (overlap with subjects in Gur <i>et al.</i> , 1991), 18–80 years old, 70 M, 60 F. No major medical, neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla). Volume measurements using a combination of computer-assisted trace methodology and pixel segmentation of 3D images reconstructed from T ₂ -weighted axial images (5 mm thick, contiguous). Good rater reliabilities, but 'blindness' not specified	Ratio of frontal lobe to IV was smaller in males over 40 years old than in younger males; no such group difference in females. In contrast, the R > L asymmetry of frontal lobe to IV was larger in older females than younger females; no such group difference in males. Ratio of temporal lobe to IV was also smaller in males over 40 years old than in younger males; no such group difference in females; no interactions with laterality. Ratio of the remaining brain volume to IV was smaller in older than younger subjects for both sexes; no interactions with laterality
DeCarli <i>et al.</i> , 1994 ⁵²	30 Healthy male volunteers, 19–92 years old. No major medical, neurologic, or psychiatric illness. 29 Right-handed	MR imaging (0.5 tesla). Volume measurements using computer-assisted trace methodology of T ₁ -weighted coronal images (6 mm thick, contiguous) through temporal lobe, by single (blind?) rater	Age associated with decreased ratio of frontal lobe volume to IV but not with temporal lobe volume to IV. No interactions with laterality
Pfefferbaum <i>et al.</i> , 1994 ⁵³	73 Healthy male volunteers (included in Pfefferbaum <i>et al.</i> , 1993), 21–70 years old (44.1 + 13.8 years). No major medical, neurologic, or psychiatric illness. Left handers included (n not specified)	MR imaging (1.5 tesla). Blinded volume measurements derived from semi-automated pixel segmentation of intermediate and T ₂ -weighted axial images ($n = 17–20$, 5 mm thick, 2.5 mm interscan gap)	Adjusting for head size, age associated with decreased cortical gray matter volume (0.7 ml/year), but not with cortical white matter volume. Interactions with laterality not reported
Soininen <i>et al.</i> , 1994 ⁵⁴	32 Healthy volunteers from the community, all with MMSE scores > 25, 16 with age-associated memory impairment (AAMI) (67.7 + 7 years; 4 M, 12 F), 16 controls (70.2 + 4.7 years; 6 M, 10 F) without AAMI. All but one right-handed	MR imaging (1.5 tesla). Blinded volume measurements using computer-assisted trace methodology of T ₁ -weighted coronal images (1 mm thick, contiguous) through temporal lobe, by single rater with established reliabilities	No group differences in hippocampal volumes, although controls (but not AAMI subjects) exhibited significant R > L asymmetry. No group differences in amygdala volume or asymmetry

continued

Table 7.2 continued

Study	Subjects	Imaging and measurement technique	Findings
Blatter <i>et al.</i> , 1995 ⁵⁵	194 Healthy volunteers, 16–65 years old, 89 M, 105 F. No history (by questionnaire) of any neurologic or psychiatric illness. 95% Right-handed	MR imaging (1.5 tesla). Volume measurements derived from semi-automated pixel segmentation and trace methodologies of intermediate and T ₂ -weighted axial images (5 mm thick, 2 mm gap). High rater reliabilities (blinded status?)	Adjusting for head size, age associated with decreased total brain volume and gray matter volume, but not white matter volume. Correlations tended to be higher for males than females, but these apparent differences were not analyzed. However, only females showed significant age-related reductions in gray matter. Interactions with laterality not reported
Convit <i>et al.</i> , 1995 ⁵⁶	37 Healthy adult volunteers, 27 older (14 M, 13 F; 69.2 + 8.3 years old), 10 younger (5 M, 5 F; 26.1 + 4.1 years old). No evidence of stroke or major medical or psychiatric illness	MR imaging (1.5 tesla). Blinded volume measurements by single rater (reliabilities?) using computer-assisted trace methods of T ₁ -weighted coronal images (4 mm thick, 10% gap)	Controlling for sex and head size, age associated with volume loss in lateral temporal lobe (especially fusiform gyrus) and medial temporal lobe (especially hippocampus and parahippocampus)
Hokama <i>et al.</i> , 1995 ⁵⁷	15 Healthy male community volunteers. 20–55 years old. No lifetime history of major medical, neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla). Volume measurements of basal ganglia using semi-automated computer assisted trace methodology from T ₁ -weighted coronal and axial sections (1.5 mm thick, contiguous) by raters with established reliabilities	Age associated with decreased volumes of caudate and putamen, but not of globus pallidus. No correlation between basal ganglia volumes and IQ as estimated by WAIS-R information subscale
Parashos <i>et al.</i> , 1995 ⁵⁸	80 Healthy volunteers (overlap with subjects in Coffey <i>et al.</i> , 1992), 30–91 years old, 28 M, 52 F. No lifetime history of neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla). Blinded area measurements using computer-assisted trace methodology of T ₁ -weighted midsagittal image (5 mm thick), made by single rater with established rater reliabilities	Adjusting for IV, increasing age associated with smaller total and regional callosal areas, especially of anterior regions (males = females)
Fox <i>et al.</i> , 1996 ⁵⁹	11 Adult volunteers with no evidence of memory impairment on testing, 5 M, 6 F; 51.3 + 5.9 years old. No additional details provided	MR imaging (1.5 tesla). Volume measurements using computer-assisted pixel segmentation of T ₁ -weighted coronal images (1.5 mm thick, contiguous). Scanning repeated at 12.8 + 4.3 months and volume differences determined from subtraction images	Over the follow-up period, brain volume decreased by 0.05% (~0.03 ml)
Janowsky <i>et al.</i> , 1996 ⁶⁰	60 Healthy elderly volunteers, 66–94 years old (mean 78.2), 15 M, 45 F. No major medical, neurologic, or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Area measurement of corpus callosum derived from computer-assisted trace methodology. Number of raters and their 'blindness' not specified	Age associated with decreased total callosal area, anterior callosal area and middle callosal area. Interactions with sex not reported
Murphy <i>et al.</i> , 1996 ⁶¹	69 Healthy volunteers. 35 M (44 + 23 years old); 34 F (50 + 21 years old). No major medical or psychiatric illness. All right-handed	MR imaging (0.5 and 1.5 tesla). Blinded volume measurements using computer-assisted segmentation and trace methodology of contiguous coronal images (5–6 mm thick). Number of raters not specified	Relative to 'young' subjects (age 20–35 years), 'old' subjects (60–85 years) had smaller brain matter volume ratios of cerebellum to IV (males = females), cerebellum to IV (males > females), frontal lobe to IV (males > females), temporal lobe to IV (males > females), parietal lobe to IV (females > males), parieto-occipital lobe to IV (males = females), parahippocampal gyrus to IV (males = females), amygdala to IV (males = females), hippocampus to IV (females > males), thalamus to IV (males = females), lenticular nucleus to IV (males = females), and caudate to IV (males = females). For the frontal lobe, the right side decreased more than the left with age in males, but in females the left side decreased more than the right. For all other regions, there were no interactions with laterality
Deshmukh <i>et al.</i> , 1997 ⁶²	10 Healthy male volunteers, 50.1 + 13.8 years old. No evidence of major medical, neurologic or psychiatric illness. Nine right-handed; one left-handed	MR imaging (1.5 tesla). Volume measures using semi-automated computer-assisted trace methodology from 3D T ₁ -weighted sagittal sections, realigned in the axial plane, by raters with established reliabilities	Age associated with decreased volume of cerebellar lobules VI–VII

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Table 7.2 continued

Study	Subjects	Imaging and measurement technique	Findings
Jack <i>et al.</i> , 1997 ⁶³	126 Healthy elderly volunteers 51–89 years old (79.15 + 6.73 years), 44 M, 82 F. No active neurologic or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Blinded volume measurements using computer-assisted trace methodology of T ₁ -weighted 3D volumetric images (1.6 mm thick, contiguous, <i>n</i> = 124) by single rater with established reliabilities	Age associated with decreased volume ratio of hippocampus to IV (45.63 ml/year), amygdala to IV (20.75 ml/year), and parahippocampal gyrus to IV (46.65 ml/year); effects similar for males and females. Effects were similar for the two hemispheres, except for the parahippocampal gyrus (L > R)
Kaye <i>et al.</i> , 1997 ⁶⁴	30 Healthy elderly volunteers from the community, with MMSE524. All 584 years old; 14 M, 16 F. No evidence of major medical, neurologic or psychiatric illness	MR imaging (1.5 tesla). Blinded volume measurements using computer-assisted trace methodology of T ₁ -weighted coronal images (4 mm thick, contiguous) by raters with established reliabilities. Scanning repeated annually over a mean of 42 months	No group differences in rate of volume loss in hippocampus (about 2%/year) or parahippocampus (about 2.5%/year)
O'Brien <i>et al.</i> , 1997 ⁶⁵	40 Healthy community volunteers, 55–96 years old, 20 M, 20 F. No evidence of major medical or neurologic illness, or of depression or drug abuse	MR imaging (0.3 tesla). Ratings of amygdala–hippocampal atrophy from T ₁ -weighted coronal images (5.1 mm thick, 0.5 mm gap) by two raters with established reliabilities, blind to cognitive scores	Age associated with presence of amygdala–hippocampal atrophy
Raz <i>et al.</i> , 1997 ⁶⁶	148 Healthy volunteers, 18–77 years old, 66 M (47.39 + 18.07 years old); 82 F (45.72 + 6.48 years old). No major medical, neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla). Blinded volume measurements (digital planimetry) from scans of T ₁ -weighted reformatted coronal images (1.3 mm thick, contiguous). Good rater reliabilities among eight raters	Adjusted for height, age significantly related to smaller volumes of whole brain (males = females), prefrontal gray matter (males = females), inferior temporal cortex (males > females), fusiform gyrus (males = females), hippocampal formation (males = females), primary somatosensory cortex (males = females), superior parietal cortex (males = females), prefrontal white matter (males = females), and superior parietal white matter (males = females). No age effects were found for anterior cingulate cortex, parahippocampal cortex, primary motor cortex, inferior parietal cortex, visual cortex, and precentral, postcentral or inferior parietal white matter. No interactions with laterality
Salat <i>et al.</i> , 1997 ⁶⁷	76 Healthy elderly volunteers, 65–95 years old (mean 77.7 years), 31 M, 45 F. No major medical or neurologic illness, and no depression. All right-handed, except one left-handed F	MR imaging (1.5 tesla). Area measurements (one of three raters with established reliabilities) of corpus callosum, pons, and cerebellum using trace methodology of T ₁ -weighted midsagittal image	Age associated with decreased total, anterior, and middle callosum areas in females but not males. No relation of age to pons or cerebellum areas
Coffey <i>et al.</i> , 1998 ⁶⁸	330 Elderly volunteers living independently in the community, 66–96 years old (74.98 + 5.09), 129 M, 201 F. No history of neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla, <i>n</i> = 248; 0.35 tesla, <i>n</i> = 82). Blinded volume measurements (one of two raters with established reliabilities) using computer-assisted trace methodology of T ₁ -weighted axial images (5 mm thick, no interscan gap)	Adjusting for IV, age associated with decreased cerebral hemisphere volume (72.79 ml/year) (males = females), frontal region area (70.13 ml/year) (males = females), temporal–parietal region area (70.13 ml/year) (males = females), and parietal–occipital region (males > females, 70.31 vs. 70.09 ml/year, respectively). All effects similar in both hemispheres
Davatzikos and Resnik 1998 ⁶⁹	114 Healthy volunteers 56–85 years old, 68 M (70.9 + 7.6 years), 46 F (69.4 + 8.0 years old). All right-handed. No additional details provided	MR imaging (1.5 tesla). Quantitative morphometry of the corpus callosum using computer-assisted trace methodology of T ₁ -weighted midsagittal image (1.5 mm thick). Morphometry was quantitated using a template and deformation function. No additional details provided	Age associated with decreased total and regional callosal size (males = females), with exception of anterior and posterior extremes

continued

Table 7.2 continued

Study	Subjects	Imaging and measurement technique	Findings
Gunning-Dixon <i>et al.</i> , 1998 ⁷⁰	Same as Raz <i>et al.</i> , 1997	MR imaging (1.5 tesla). Blinded volume measurements (digital planimetry) from scans of T ₁ -weighted reformatted coronal images (1.3 mm thick, contiguous). Good rater reliabilities among eight raters	Age associated with decreased caudate volume (L > R in males, R > L in females), decreased putamen volume (R > L, males = females), and decreased globus pallidus volume (males only)
Gur <i>et al.</i> , 1998 ⁷¹	17 Healthy volunteers (overlap with subjects in Gur <i>et al.</i> , 1991 and Cowell <i>et al.</i> , 1994) 31.9 + 8.9 years old, 13 M, 4 F	MR imaging (1.5 tesla). Volume measurements using a combination of computer-assisted trace methodology and pixel segmentation of 3D images reconstructed from T ₂ -weighted axial images (5 mm thick, contiguous). Good rater reliabilities, but 'blindness' not specified. Scanning repeated an average of 32 months later.	No significant change over follow-up period in whole brain, CSF or frontal lobe volumes. Significant volume loss was observed for L (7.5%) and R (7.2%) temporal lobes
Guttmann <i>et al.</i> , 1998 ⁷²	72 Healthy volunteers, 18–81 years old, 22 M, 50 F. No history of psychiatric illness, epilepsy, or severe head trauma. Handedness not specified	MR imaging (1.5 tesla). Blinded volume measurements using computer-assisted segmentation and trace methodology of contiguous axial images (3 mm thick). Good interrater reliabilities	Age associated with decreased ratio of total white matter volume to IV and decreased total gray matter volume to IV. Interactions with sex or laterality not reported
Jack <i>et al.</i> , 1998 ⁷³	24 Elderly volunteers, 70–89 years old (81.04 + 3.78 years), 8 M, 16 F. No active neurologic or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Blinded volume measurements using computer-assisted trace methodology of T ₁ -weighted 3D volumetric images (1.6 mm thick, contiguous, <i>n</i> = 124) by single rater with established reliabilities	Over a 12 month interval, mean hippocampal volume decreased by 1.55% and mean temporal horn volume increased by 6.15% (males = females, L = R)
Laakso <i>et al.</i> , 1998 ⁷⁴	42 cognitively normal healthy elderly community volunteers, 64–79 (72 + 4) years old, 19 M, 23 F	MR imaging (1.5 tesla). Volume measures using computer-assisted trace methods from T ₁ -weighted coronal images (~2 mm thick, contiguous) by a single blinded rater with established reliabilities	No relation between age and hippocampal atrophy
Mueller <i>et al.</i> , 1998 ⁷⁵	46 Healthy elderly volunteers 65–74 years old (6 M, 5 F), 75–84 years old (8 M, 7 F) and 85–95 years old (9 M, 11 F). All functionally independent, MMSE524, and free of major medical and neurologic illness, as well as depression. Handedness not specified	MR imaging (1.5 tesla). Volume measurements (non-blind?) using computer-assisted pixel segmentation of contiguous coronal images (4 mm thick). Excellent interrater reliabilities. Scanning repeated annually or biannually over 3–9 year follow-up	Adjusting for IV, age associated with decreased volumes of total brain, hemispheres, frontal lobes, temporal lobes, basilar-subcortical region, hippocampus, and hippocampal gyrus. Interactions with sex not reported. Over the follow-up period, significant volume decreases were seen in hippocampus (~0.02 ml/year), parahippocampal gyrus (in youngest group only, ~0.05 ml/year), parietal-occipital region (in middle and oldest age groups, ~3 ml/year), and basilar region (in middle group only, ~0.5 ml/year). No volume decreases were seen in hemispheres, frontal lobes or temporal lobes
Oguro <i>et al.</i> , 1998 ⁷⁶	152 Healthy adults, 81 M, 71 F, age range 40s–70s. No evidence of neurological disease	MR imaging (0.2 tesla). Linear and area measurements using computer-assisted trace methodology from T ₁ -weighted midsagittal image (7 mm thick) and T ₂ -weighted axial images (no details), by raters (<i>n</i> = ?) with established reliabilities	Age associated with decreased linear measures of midbrain tegmentum (males only), midbrain prectectum (males and females), and base of pons (males only), but not with pontine tegmentum or fourth ventricle. Age associated with decreased area of cerebellar vermis (males only), but not of pons. Age associated with decreased ratio of cerebrum to IA (males and females) at level of third ventricle and at level of body of lateral ventricles. Interactions with laterality not reported
Pfefferbaum <i>et al.</i> , 1998 ⁷⁷	28 Healthy male volunteers (overlap with Pfefferbaum <i>et al.</i> , 1994), 21–68 years old (51 + 13.8 years). No major medical, neurologic, or psychiatric illness. Left-handers included (<i>n</i> not specified)	MR imaging (1.5 tesla). Blinded volume measurements derived from semi-automated pixel segmentation of intermediate and T ₂ -weighted axial images (<i>n</i> = 17–20, 5 mm thick, 2.5 mm interscan gap). Scanning repeated at 5 year follow-up	Over the follow-up interval, significant decrease in total gray matter volume and in regional gray matter volume (pre-frontal gray ~2 ml, or 7%; posterior parieto-occipital gray ~1 ml, or 3.5%) (no change in frontal, anterior superior temporal, posterior superior temporal, or anterior parietal regions gray matter volume). Interactions with laterality not reported
Raz <i>et al.</i> , 1998 ⁷⁸	146 Healthy volunteers (overlap with Raz <i>et al.</i> , 1997), 18–77 years old, 64 M (48 + 18 years); 82 F (46 + 17 years). No evidence of major medical, neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla). Blinded volume measurements (digital planimetry) from scans of T ₁ -weighted reformatted coronal and sagittal images (0.8 mm thick, 1.5 mm thick). Good rater reliabilities	Age associated with volume loss in cerebellar hemispheres (~2%/decade), vermis, vermian lobules VI and VII (~4%/decade), and posterior vermis (lobules VIII-X; ~2%/decade), but not in anterior vermis (lobules I–V) or pons

Table 7.3. Aging and changes in size of CSF spaces

Study	Subjects	Imaging and measurement technique	Findings
Barron <i>et al.</i> , 1976 ⁷	135 Volunteers, 9 months–90 years old, equal gender distribution in all age groups (8 M, 7 F per decade). No history of neurological disease; psychiatric history not reported. Handedness not specified	Computed tomography (CT). Planimetric determination of ventricular–brain ratio (VBR) by single rater (average of three measurements) from Polaroid photograph	Age associated with increased VBR and with increased variability in VBR. Interactions with sex or laterality not reported
Earnest <i>et al.</i> , 1979 ⁷⁹	59 Volunteer retirees, 60–99 years old, 11 M, 48 F. Living independently and free of neurological disease. Handedness not specified	CT. Linear and planimetric measures of ventricular size at three different levels, from photographs. Linear measurements of four largest sulci. No additional data provided	Subjects 80 years or older ($n=29$) had larger ratio of ventricular size to intracranial size than did younger subjects ($n=30$). The sum of the widths of the four sulci was greater in older subjects than in younger subjects. Interactions with sex or laterality not reported
Jacoby <i>et al.</i> , 1980 ¹⁹	50 Healthy elderly volunteers, 62–88 years old, 10 M, 40 F. No history of significant psychiatric or neurologic illness. Handedness not specified	CT. Ratings (small, normal, enlarged) of ventricular size from films by single blinded rater (rater reliability not reported). Planimetric determination of ventricular–skull ratio and Evans's ratio from films by single rater (average of three measurements) with established reliabilities	8 (16%) Subjects were rated as having 'enlarged' lateral ventricles. No significant correlation between age and ventricular–skull ratio or Evans's ratio. Interactions with sex or laterality not reported. Adjusting for age, no relation between any CT measure and performance on the Hodkinson test of memory and orientation
Meese <i>et al.</i> , 1980 ⁸⁰	160 Healthy 'volunteers', 1–71 years old, 10 M and 10 F in each decade. No additional data provided	CT. Linear measurements of ventricular size and sulcal width from four axial slices (no additional data provided)	Apparent age-related changes in some measures of ventricular size and sulcal width, but these changes not analyzed statistically. Interactions with sex or laterality not reported
Cala <i>et al.</i> , 1981 ²⁰	115 Volunteers, 15–40 years old, 62 M, 53 F. No history of migraine, head trauma, or excessive alcohol intake (no additional details provided). All but eight subjects right-handed	CT. (n = two scanners). Planimetric measurements of ventricular–skull ratio at level of frontal horns (no additional details provided). Axial slices (13 mm thick)	No relationship between age and ventricular–skull ratio. Interactions with sex or laterality not reported
Soiminen <i>et al.</i> , 1982 ⁸¹	85 Volunteers, 53 from community and 32 from nursing home, 75+7 years old, 23 M, 62 F. No neurological disease (no additional details provided)	CT. Linear measurements (from films?) of ventricular and sulcal size. Axial slices ($n=8-12$, 8 mm thick). No additional details provided	Age correlated with ratios of ventricular width to skull width (frontal horn index and cella media index). Age correlated with mean width of four largest sulci. Correlations were found between a composite neuro-psychological test score and the size of the lateral and IIIrd ventricles, the left Sylvian fissure, and the right temporal horn, but the effects of age were not controlled. Interactions with sex or laterality not reported
Zatz <i>et al.</i> , 1982a ¹²	123 Volunteers, 10–90 years old, 49 M, 74 F. No history of neurological or major medical disease. Handedness not specified	CT. Volume measurement derived from computer-assisted pixel segmentation technique (ASI-II program). Axial slices ($n=9$, 10 mm thick, 10 mm interscan gap)	Age significantly associated with increased ventricular volume (males = females), even after controlling for intracranial volume (IV). Increased variability of ventricular size with age. Age associated with increased sulcal CSF volume, even after controlling for IV
Gado <i>et al.</i> , 1983 ¹⁴	12 Elderly volunteers, 64–81 years old, 9 M, 3 F. No additional clinical data provided	CT. Volume measurements derived from computer-assisted pixel segmentation technique (seventh axial slices, 8 mm thick). Linear measurements from axial images. Number of raters and rater reliabilities not specified	During 1 year follow-up, ratio of ventricular volume to IV increased significantly by an average of 3.7%. No significant changes in linear measures of ventricular size (VBR, IIIrd ventricular ratio, frontal horn ratio). During 1 year follow-up, ratio of sulcal volume to IV increased significantly by an average of 13%. Interactions with sex or laterality not reported
Laffey <i>et al.</i> , 1984 ¹¹	212 Elderly volunteers, 65–89 years old, 110 M; 102 F. No evidence of alcoholism, dementia, or neurologic illness	CT. Qualitative rating (six-point scale) of ventricular enlargement and sulcal widening from films, by two experienced radiologists with established reliabilities	Age associated with increased ventricular size. No association between age and ratings of sulcal widening. Interactions with sex or laterality not reported

continued

Table 7.3 continued

Study	Subjects	Imaging and measurement technique	Findings
Gomori <i>et al.</i> , 1984 ²¹	148 Neurologically intact volunteers, 28–84 years old. Subjects with minor neurological symptoms, patients with lung, breast, prostate, colon cancer but no CNS involvement. Neurologic exam normal	CT. Linear (sum of bicaudate and Sylvian fissure ratios)	CSF space increased with age; age correlated with the sum of the two ratios (bicaudate and Sylvian fissure, $r=0.64$). The sum of the ratios increased two-fold between ages 28–49 and 80–89, faster after the fifth decade
Takeda <i>et al.</i> , 1984 ⁸²	980 Patients scheduled for CT, 10–88 years old, 483 M, 497 F. Included patients with hypertension, diabetes mellitus, ischemic heart disease, lung disease and renal disease. Excluded subjects with abnormal CT	CT. Volumetric. Axial slices ($n=7$) starting at the supraorbitomeatal line (10 mm thick, contiguous sections)	CSF space increased with age; CSF space volume displayed five-fold increase from age 40 (21.0 cm ³) to 90 (124.3 cm ³). Brain atrophy index (BAI) showed similar increase. Both CSF space volume and BAI were stable before age 40. Brain atrophy was accelerated in males during the fourth decade and the rate decreased afterwards, while atrophy in females proceeded with a stable rate throughout the lifespan. Increased variability with age
Schwartz <i>et al.</i> , 1985 ⁵	30 Healthy male volunteers, 21–81 years old. No history of major medical, neurologic, or psychiatric illness. Handedness not specified	CT. Volume measurements derived from computer-assisted pixel segmentation technique (ASI-II program). Axial slices ($n=7$) starting from the plane of the inferior orbito-meatal line (10 mm thick, 7 mm interscan gap)	Age correlated with areas and volumes of lateral and IIIrd ventricles, even after adjusting for height and intracranial area. Age correlated with VBR. Increased variability of ventricular size with age. No laterality effects. Age correlated with CSF volume (ventricular plus basal cisterns), even after controlling for IV. Increased variability of CSF volume with age
Pfefferbaum <i>et al.</i> , 1986 ¹³	57 Healthy volunteers, 20–84 years old, 27 M, 30 F. No additional data provided	CT. Volume measurements derived from computer-assisted pixel segmentation technique (modification of Gado <i>et al.</i> , 1983). Contiguous axial slices ($n=5$), starting at the level of the superior roof of the orbits	Age associated with increased ratio of ventricular volume to IV. Increased variability in ventricular volume with age. Interactions with sex or laterality not reported. Age associated with increased ratio of sulcal CSF volume to IV (from single axial slice [8 mm thick] approximately 48 mm from the level of the superior roof of the orbits). Age associated with increased variability of sulcal CSF volume to IV
Nagata, <i>et al.</i> , 1987 ⁸³	500 Patients with head CT, 10–90 years old. Excluded: subjects with abnormal CT	CT. Volumetric. Axial slices ($n=6$), beginning at the level of the basal cistern (10 mm thick)	CSF space increased with age; four-fold increase in BAI between ages 50 (2.3%) and 80 (10%). BAI was stable before the sixth decade. Increased variability with age
Stafford <i>et al.</i> , 1988 ²²	79 Healthy male volunteers, 31–87 years old. No severe medical or psychiatric illness. Handedness not specified	CT. Volume measurements derived from computer-assisted pixel segmentation technique (ASI-II program). Axial slices ($n=3$) at mid-, high- and supraventricular levels	Age associated with increased ratio of ventricular–brain volume. Age associated with increased ratio of supraventricular CSF–brain volume. Interactions with laterality not reported. Inverse correlation observed between a discriminant function of ventricular volume measures and a discriminant function of neuropsychological tests of naming and abstraction
Pearlson <i>et al.</i> , 1989 ⁸⁴	31 Healthy elderly volunteers, 68.3 + 1.2 years old, 15 M, 16 F. No major medical, neurologic, or psychiatric illness. Handedness not specified	CT. Planimetric determination of VBR from films by one of two raters, each with established reliabilities	Age correlated with VBR. Interactions with sex or laterality not reported
Kaye <i>et al.</i> , 1992 ⁸⁵	107 Healthy volunteers, 64 M (21–90 years old); 43 F (23–88 years old). No major medical, neurologic, or psychiatric illness. Handedness not specified	CT. Volume measurements derived from computer-assisted pixel segmentation technique (ASI-II program). Axial slices (10 mm thick, 7 mm interscan gap)	Age associated with increased ventricular volume in both males and females (about 20% per decade); precipitous increases observed beginning in the fifth decade in males and in the sixth decade in females. Interactions with laterality not reported

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Table 7.3. *continued*

Study	Subjects	Imaging and measurement technique	Findings
Sullivan <i>et al.</i> , 1993 ⁸⁶	114 Healthy volunteers, 21–82 years old (51.2 + 17.7 years), 84 M, 30 F. No history of major medical, neurologic, or psychiatric illness. 90% Right-handed	CT. Volume measurements derived from computer-assisted pixel segmentation technique (modification of Gado <i>et al.</i> , 1983). Axial slices ($n = 10$, 10 mm thick). 10 neuropsychological tests (MMSE, Trail Making Test A and B, WAIS-R subtests—Information, Digit Span, Vocabulary, Digit Symbol, Picture Completion, Block Design, and Object Assembly)	Age correlated with total and third ventricular volume, even after adjustments for head size (males = females). No correlation between age related changes in total or third ventricular volume and performance on 10 neuropsychological tests. Age correlated with increased CSF volume in Sylvian fissure and in vertex, frontal and parieto-occipital sulci (males = females).
Shear <i>et al.</i> , 1995 ⁸⁷	35 Healthy volunteers (included in Sullivan <i>et al.</i> , 1993), 67.4 + 7.4 years old, 23 M, 12 F. No history of major medical, neurologic, or psychiatric illness	CT. Longitudinal within-subject follow-up, using blinded volume measurements per technique of Sullivan <i>et al.</i> , 1993. High rater reliabilities	Over mean (+SD) follow-up of 2.6 (+0.96) years, increases were observed in CSF volumes of frontal sulci (0.31 ml/year), Sylvian fissure (0.58 ml/year), parieto-occipital sulci (0.05 ml/year), and ventricular system (0.61 ml/year). Interactions with sex not reported
Elwan <i>et al.</i> , 1996 ⁸⁸	88 Healthy 'lower middle class' volunteers, 40–76 years old (54.8 + 9.6 years), 57 M, 31 F. No major medical, neurologic, or psychiatric illness. All right-handed	CT. Multiple distance measurements (no additional details provided)	Age correlated with maximum width of IIIrd ventricle. Interactions with sex or laterality not reported
Grant <i>et al.</i> , 1987 ⁸⁹	64 Healthy volunteers, 18–64 years old, 25 M, 39 F. No history of neurological disease; psychiatric history not reported. Handedness not specified	Magnetic resonance (MR) imaging (0.15 tesla). Mathematically derived estimate of ventricular volume and CSF volume from signal intensity measurements made on single sagittal slice (number of raters not specified)	Age associated with increased ventricular volume in males, but not females; however, this apparent gender difference was not tested statistically. Interactions with laterality not reported. Age associated with increased total (ventricular plus cisternal) cranial CSF volume (males = females). No control for size of brain or head
Condon <i>et al.</i> , 1988 ²⁹	40 Volunteers, 20–60 years old, 20 M, 20 F. No additional details provided	MR imaging (0.15 tesla). Volume measurement (two raters) derived from computer-assisted pixel segmentation of contiguous sagittal slices (variable slice thickness and number)	For males but not females, age correlated with ratio of total ventricular volume to IV, total CSF volume to IV, and total sulcal CSF volume to IV
Yoshii <i>et al.</i> , 1988 ³⁰	58 Healthy volunteers, 21–81 years old, 29 M, 29 F. Neurologic and psychiatric histories not reported. Handedness not specified	MR imaging (1.0 tesla). Blinded global ratings (four-point scale) of lateral ventricular enlargement from inversion recovery films (axial slices, n unspecified, 10 mm thick, 3 mm interscan gap). Numbers of raters and rater reliabilities not specified	Age correlated with ratings of lateral ventricular enlargement (males = females). Interactions with laterality not reported
Jernigan <i>et al.</i> , 1990 ³³	58 Healthy volunteers, 8–79 years old, 35 M, 23 F. No neurologic, psychiatric, or medical (e.g. diabetes mellitus and heart disease) illness. Handedness not specified	MR imaging (1.5 tesla). Volume estimates (one of two raters) derived from computer-assisted pixel classification of multiple spin-echo axial images (5 mm thick, 2.5 mm interscan gap)	Age associated with increased ratio of ventricular CSF volume to IV. Age associated with increased ratio of sulcal CSF volume to IV. Interactions with sex or laterality not reported
Wahlund <i>et al.</i> , 1990 ⁹⁰	24 Healthy elderly volunteers, 75–85 years old (mean = 79 years), 8 M, 16 F. No evidence of neurologic or psychiatric illness. Handedness not specified	MR imaging (0.02 tesla). Visual ratings (5-point scale) of CSF spaces on T ₂ -weighted axial films (slice = 10 mm thick, no gap) by 2 raters (blind?) with established reliabilities. Area measurements based upon computer-assisted pixel classification technique, from single axial section at level of basal ganglia	No correlation between age and visual ratings or area measurements of sulcal CSF or lateral ventricle CSF size
Gur <i>et al.</i> , 1991 ³⁷	69 Healthy volunteers, 18–80 years old, 34 M, 35 F. No neurologic or psychiatric illness. 66 Right-handed; three left-handed	MR imaging (1.5 tesla). Volume measurements (any two of four raters) derived from segmentation technique based on two-feature pixel classification of multiple spin-echo axial images (5 mm thick, contiguous)	Older (555 years) subjects ($n = 26$) had larger total CSF volume (males > females), larger ratio of ventricular CSF volume to IV (males = females), and larger ratio of sulcal CSF volume to IV (males > females). Effects of age on ratio of ventricular CSF volume to IV were asymmetric (L > R) in males but not in females

continued

Table 7.3 continued

Study	Subjects	Imaging and measurement technique	Findings
Tanna <i>et al.</i> , 1991 ⁴⁰	16 Healthy volunteers, 52–86 years old, 5 M, 11 F. No evidence of major medical, neurologic or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Volume measurements (one of two raters with established reliabilities) derived from segmentation techniques based on two-feature pixel classification of multiple spin-echo axial images (5 mm thick, 2.5 mm interscan gap)	Age significantly correlated with ratio of ventricular CSF volume to total CSF plus total brain volume. Trend (non-significant) for age to be associated with increasing ratio of sulcal CSF volume to total CSF plus total brain volume. Interactions with sex or laterality not reported
Coffey <i>et al.</i> , 1992 ⁴¹	76 Healthy volunteers, 36–91 years old, 25 M, 51 F. No lifetime evidence of neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla). Volume measurements (one of three blinded raters with established reliabilities) using computer-assisted trace methodology of T ₁ -weighted coronal images ($n = 30$ – 35 , 5 mm thick, contiguous). Blinded clinical ratings (five-point scale) of lateral ventricular enlargement from films (average score of two experienced raters)	Adjusting for IV, age associated with increased volumes of the third (2.8%/year) and lateral (3.2%/year) ventricles (males = females). Age associated with increased odds (7.7%/year) of at least mild lateral ventricular enlargement, from 0.10 at age 40 to 2.22 at age 80 (males = females). No interactions with laterality
Lim <i>et al.</i> , 1992 ⁴⁴	14 Healthy male volunteers, 8 young (21–25 years old), 6 elderly (68–76 years old). No evidence of significant medical or psychiatric illness. Handedness not specified	MR imaging (1.5 tesla). Blinded volume measurements derived from semi-automated pixel segmentation of intermediate and T ₂ -weighted axial imaging ($n = 8$, 5 mm thick, 2.5 mm interscan gap)	Compared with younger males, older males had higher percentage of CSF volume to IV (8% vs. 20.1%)
Matsubayashi <i>et al.</i> , 1992 ⁹¹	73 Healthy volunteers, 59–83 years old, 24 M, 49 F. No history of major medical, neurologic, or psychiatric illness	MR imaging (0.5 tesla). Planimetric determination of ventricular–parenchymal ratio (VPR). No additional details provided	Age correlated with VPR. Interactions with sex or laterality not reported
Murphy <i>et al.</i> , 1992 ⁴⁵	27 Healthy males, 19–92 years old. No major medical, neurologic, or psychiatric illness. Handedness not specified	MR imaging (0.5 tesla). Blinded volume measurements derived from semi-automated pixel segmentation of proton density axial images ($n = 36$, 7 mm thick, contiguous). Rater reliabilities established, but number of raters not specified	Compared with younger males (under 60 years old; $n = 10$), older males ($n = 17$) had larger ratios of lateral ventricular volume to IV, larger ratio of third ventricular volume to IV, and larger ratios of peripheral CSF volume (total CSF volume minus ventricular volumes) to IV. No interactions with laterality
Raz <i>et al.</i> , 1993a ⁴⁷	29 Healthy volunteers, 18–78 years old, 17 M, 12 F. No major medical, neurologic, or psychiatric illness. Self-reported right-handers	MR imaging (0.30 tesla). Blinded volume measurements from films using digital planimetry of T ₁ -weighted and proton density sagittal and coronal images. Good rater ($n = 2$) reliabilities	Controlling for head size, age associated with increased lateral ventricular volume (males = females). Interactions with laterality not reported
Christiansen <i>et al.</i> , 1994 ⁵⁰	142 Healthy volunteers, 21–80 years old, 78 M, 64 F. No major medical or neurologic illness	MR imaging (1.5 tesla). Volume measurements using manual tracing of T ₂ -weighted axial images (4 mm thick, 4 mm interscan gap). No additional details provided	Age associated with increased lateral ventricle volume in males (134%) and females (66%), but these apparent gender differences were not statistically compared. Interactions with laterality not reported
DeCarli <i>et al.</i> , 1994 ⁵²	30 Healthy male volunteers, 18–92 years old. No major medical, neurologic, or psychiatric illness. 29 Right-handed	MR imaging (0.5 tesla). Volume measurements of T ₁ -weighted coronal images (6 mm thick, contiguous) by single rater using computer-assisted pixel segmentation techniques. Good rater reliabilities	Age associated with increased volume of sulcal CSF to IV (1.3%/decade), central CSF to IV (0.3%/decade), and third ventricle CSF to IV (0.04%/decade). For all measures, no interactions with laterality
Pfefferbaum <i>et al.</i> , 1994 ⁵³	73 Healthy male volunteers (included in Pfefferbaum <i>et al.</i> , 1993), 21–70 years old (44.1 + 13.8 years). No major medical, neurologic, or psychiatric illness. Left-handers included (n not specified)	MR imaging (1.5 tesla). Blinded volume measurements of T ₂ -weighted axial slices (5 mm thick, 2.5 mm interscan gap) by four raters using computer-assisted pixel segmentation techniques. Good rater reliabilities	Age associated with increased cortical CSF volume to IV (0.6 ml/year) and ventricular volume to IV (0.3 ml/year). Interactions with laterality not reported
Blatter <i>et al.</i> , 1995 ⁵⁵	194 Healthy volunteers, 16–65 years old, 89 M, 105 F. No history (by questionnaire) of any neurologic or psychiatric illness. 95% Right-handed	MR imaging (1.5 tesla). Volume measurements derived from semi-automated pixel segmentation and trace methodologies, of intermediate and T ₂ -weighted axial images (5 mm thick, 2 mm gap). High rater reliabilities (blinded status?)	Adjusting for IV, age associated with increased subarachnoid CSF volume, and lateral and IIIrd ventricular volumes, but not IVth ventricular volume; correlations tended to be higher for males than females, but these apparent differences were not analyzed. Interactions with laterality not reported

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Table 7.3 continued

Study	Subjects	Imaging and measurement technique	Findings
Murphy <i>et al.</i> , 1996 ⁶¹	69 Healthy volunteers, 35 males (mean + SD age = 44 + 23 years); 34 females (mean + SD age = 50 + 21 years). No major medical or psychiatric illness. All right-handed	MR imaging (0.5 and 1.5 tesla). Blinded volume measurements using computer-assisted segmentation and trace methodology of contiguous coronal images (5–6 mm thick). Number of raters not specified	Relative to younger subjects (age 20–35 years), older subjects (60–85 years) had larger ratios of lateral ventricular volume to IV (males = females), third ventricular volume to IV (females > males), and peripheral CSF volume to IV (males = females). No interactions with laterality
Salonen <i>et al.</i> , 1997 ⁹²	61 Healthy volunteers, 30–86 years old, 30 M, 31 F. No neurological symptoms or disease; psychiatric history not specified. Handedness not specified	MR imaging (1.0 tesla). Qualitative ratings (five-point scale) of sulcal and lateral ventricular enlargement. Linear measurement of maximum width of third ventricle. T ₁ -weighted axial slices (5 mm thick, 1 mm interscan gap). Number of raters and rater reliabilities not specified	Age associated with increased ratings of sulcal widening and lateral ventricular enlargement, and with width of IIIrd ventricle. Interactions with sex or laterality not reported
Yue <i>et al.</i> , 1997 ⁹³	1488 Healthy elderly volunteers from the Cardiovascular Health Study, 65–80+ years old, numbers of males and females not specified. Handedness not specified. No major medical or neurologic illness (psychiatric illness not assessed)	MR imaging (0.35 or 1.5 tesla). Blinded ratings of sulcal prominence (10-point scale) and ventricular size (10-point scale) from T ₁ -weighted axial images. Good to excellent rater reliabilities, but number of raters not specified	Age associated with increased sulcal prominence and ventricular enlargement (males = females)
Coffey <i>et al.</i> , 1998 ⁶⁸	330 Elderly volunteers living independently in the community, 66–96 years old (74.98 + 5.09), 129 M, 201 F. No history of neurologic or psychiatric illness. All right-handed	MR imaging (1.5 tesla, <i>n</i> = 248; 0.35 tesla, <i>n</i> = 82). Blinded volume measurements (one of two raters with established reliabilities) using computer-assisted trace methodology of T ₁ -weighted axial images (5 mm thick, no interscan gap)	Adjusting for IV, age associated with increased peripheral (sulcal) CSF volume, lateral fissure CSF volume, lateral ventricular volume (0.95 ml/year), and IIIrd ventricular volume (0.05 ml/year). Males showed greater age-related changes than females for peripheral CSF (2.11 ml/year vs. 0.06 ml/year, respectively) and lateral fissure volumes (0.23 ml/year vs. 0.10 ml/year, respectively). No interactions with laterality
Guttmann <i>et al.</i> , 1998 ⁷²	72 Healthy volunteers, 18–81 years old, 22 M, 50 F. No history of psychiatric illness, epilepsy, or severe head trauma. Handedness not specified	MR imaging (1.5 tesla). Blinded volume measurements using computer-assisted segmentation and trace methodology of contiguous axial images (3 mm thick). Good interrater reliabilities	Age associated with increased ratio of total CSF volume to IV. Interactions with sex or laterality not reported
Mueller <i>et al.</i> , 1998 ⁷⁵	46 Healthy elderly volunteers. 65–74 years old (6 M, 5 F), 75–84 years old (8 M, 7 F), and 85–95 years old (9 M, 11 F). All functionally independent, MMSE524, and free of major medical and neurologic illness, as well as depression. Handedness not specified	MR imaging (1.5 tesla). Volume measurements (non-blind?) using computer-assisted pixel segmentation of contiguous coronal images (4 mm thick). Excellent interrater reliabilities. Scanning repeated annually or biannually over 3–9 year follow-up	Adjusting for IV, age associated with increased temporal horn volume, but not with total CSF, sulcal CSF, or lateral ventricle volumes. Interactions with sex not reported. Over the follow-up period, significant increases were seen only in total CSF volume (~1.5 ml/year, females > males) and in lateral ventricular volume (~1.4 ml/year) (males = females)
Pfefferbaum <i>et al.</i> , 1998 ⁷⁷	28 Healthy male volunteers (overlap with Pfefferbaum <i>et al.</i> , 1994) 21–68 years old (51 + 13.8 years). No major medical, neurologic, or psychiatric illness. Left-handers included (<i>n</i> not specified)	MR imaging (1.5 tesla). Blinded volume measurements derived from semi-automated pixel segmentation of intermediate and T ₂ -weighted axial images (<i>n</i> = 17–20, 5 mm thick, 2.5 mm interscan gap). Scanning repeated at 5 year follow-up	Over the follow-up interval, significant increase in volume of lateral (~5 ml, or 20%) and third ventricles, but not of cortical sulcal CSF volume

17% over the first nine decades of life. Studies using linear measurements show a two-fold enlargement while volumetric studies report a four- to five-fold volume increase. The changes in the size of CSF spaces are accelerated after midlife (60 years). The variance in measures of CSF spaces also increases with age.

The majority of the studies report a progressive enlargement of the lateral and third ventricles with age, which ranges from less

than one to three times their initial sizes. No relationship has been found thus far between aging and fourth ventricular size. As expected, studies using linear measurements show the smallest changes, while studies using planimetric and volumetric measurements agree in finding relatively greater enlargement. The increase in ventricular size is relatively minimal until the sixth decade, after which it becomes more pronounced. Finally, the variability of the measurements of ventricular size increases with age.

Table 7.4. Aging and incidence of subcortical hyperintensities (SH)

Study	Subjects	Imaging and measurement technique	Findings
George <i>et al.</i> , 1986 ⁹⁴	47 Normal volunteers; two age groups: < 45 years (<i>n</i> = 35) and 46–78 years (<i>n</i> = 12)	MR imaging (0.3 tesla). Qualitative rating (5-point scale) of subcortical hyperintensities (white and gray matter). Axial slices (<i>n</i> = 7) 8 mm thick with 1.2 cm interscan gap	Subcortical hyperintensities increased with age; no subject under 45 had foci of increased signal; 8/12 subjects over 46 showed single or multiple foci of increased signal
Fazekas, 1989 ⁹⁵	87 Normal volunteers, 31–83 years old, 40 M, 27 F. Excluded patients with cardiovascular or cerebrovascular disease, hypertension, hyperglycemia, and neurologic or psychiatric illness	MRI (1.5 tesla). Qualitative rating (four-point scale) of subcortical hyperintensities. Axial slices (<i>n</i> = 7) intermediate and T ₂ -weighted, 5–8 mm thick	The incidence and severity of subcortical hyperintensities increased with age and in the presence of risk factors for vascular disease
Hendrie <i>et al.</i> , 1989 ⁹⁶	27 normal volunteers, 63–86 years old, 10 M, 17 F	MRI (1.5 tesla). Consensus rating (four-point scale) by two blinded raters; axial slices (T ₂ -weighted) 10 mm thick	Of the population 56.7% had SH; mean age increased with severity of SH
Jernigan <i>et al.</i> , 1991 ³³	58 Normal volunteers, 8–79 years of age, 35 M, 23 F. Excluded: diabetes mellitus, heart disease, substance abuse, developmental intellectual abnormality, psychiatric illness	MRI (1.5 tesla). Volumetric (ratio of volume of white and grey matter subcortical hyperintensities divided by cranial volume, as derived from computer-assisted pixel classification system). Axial slices of entire brain with thickness 5 mm and interscan gap 2.5 mm	SH volume ratio increased curvilinearly with age (especially after 55 years of age) from slightly over 0% at age 55 to 25% of the supratentorial volume at age 80. Increased variability with age
Matsubayashi <i>et al.</i> , 1992 ⁹¹	73 Healthy volunteers, 59–83 years of age, 24 M, 49 F. No major medical, neurological or psychiatric illness	MRI (0.5 tesla). Rating (four-point scale) of periventricular SH on T ₂ -weighted axial slices (no additional details reported)	Subjects with highest SH rating (<i>n</i> = 19) significantly older than other groups
Boone <i>et al.</i> , 1992 ⁹⁷	100 Healthy volunteers, 45–83 years of age, 36 M, 64 F. No major medical, neurological or psychiatric illness	MRI (1.5 tesla). Computer-assisted area measurements of SH from T ₂ -weighted axial sections by single rater (additional rater information not reported)	Age greater in those subjects with the largest lesion areas
Almkvist <i>et al.</i> , 1992 ⁹⁸	23 Healthy volunteers, 75 years of age and older, 9 M, 14 F. No major medical, neurological, or psychiatric illness	MRI (0.02 tesla). Area measurements of SH from T ₂ -weighted axial sections by single blinded rater (reliabilities not reported)	No correlation between age and SH area
Coffey <i>et al.</i> , 1992 ⁴¹	76 Healthy volunteers, 36–91 years old, 25 M, 51 F. No history of neurological or psychiatric illness. All right-handed	MRI (1.5 tesla). Consensus ratings of SH from intermediate and T ₂ -weighted axial films (four-point scale) by two blinded raters with established reliabilities	Increasing age associated with increased odds of SH in the deep white matter (6.3%/year) and pons (8.1%/year)

Table 7.4 summarizes the effects of aging on the incidence on MRI of foci of T₂-signal hyperintensity in the subcortical white matter and gray matter nuclei ('subcortical hyperintensities'). While the studies are difficult to compare because of differences in the definition/rating of subcortical hyperintensity, it is clear that the changes increase with age as well as in the presence of risk factors for vascular disease (e.g. smoking, hypertension, diabetes mellitus, coronary/peripheral vascular disease). In our study of 75 healthy adults⁴¹, subcortical hyperintensity was present in the deep white matter in 64.0% of subjects, in the periventricular white matter in 12.0%, in the basal ganglia in 12% and in the pons in 21.3%. The odds of having subcortical hyperintensity increased from 5% to 9% per year, depending on the anatomic region involved.

QUANTITATIVE MRI AT THE HENRY FORD AGING PROGRAM

One specific aim of our Neuropsychiatry Program at Duke University Medical Center has been to examine brain structure and function in normal aging and in patients with various psychiatric illnesses, particularly affective disorders^{41,99–102}. All participants are strongly right-handed and each receives an extensive medical and neuropsychiatric history interview and

examination. The normal subjects have no history or clinical evidence of any psychiatric disorder or any illness referable to the brain. The majority of these 'normal' subjects are also free of medical illness, so that our cohort is felt to reflect primarily 'successful' rather than 'usual' aging³⁸.

The MRI scans of the brain are performed on high-field strength (1.5 tesla) systems. Spin-echo pulse sequences are used to produce T₁-weighted (TR = 500 ms, TE = 20 ms), intermediate (TR = 2500 ms, TE = 40 ms) and T₂-weighted (TR = 2500 ms, TE = 80 ms) brain images. Slices are interleaved, relatively thin (1–5 mm) and cover the entire extent of the brain in all imaging planes. Structures of interest are outlined using computer-assisted edge detection and trace methodology (Figure 7.1). The area (cm²) within the outline is calculated automatically and volume (ml) can be determined by multiplying the area by the slice thickness and summing over the multiple slices in which the structure appears^{29,38}.

The intermediate and T₂-weighted MRI images are obtained because they are more sensitive to some forms of pathologic tissue than the T₁-weighted scans and because they permit an assessment of subcortical hyperintensities. Both the intermediate and T₂-weighted scans are acquired in the axial plane of orientation. The hard copies of these scans are coded and randomized, intermixed with scans from other patient populations, and then formally rated for subcortical hyperintensities²⁶ by

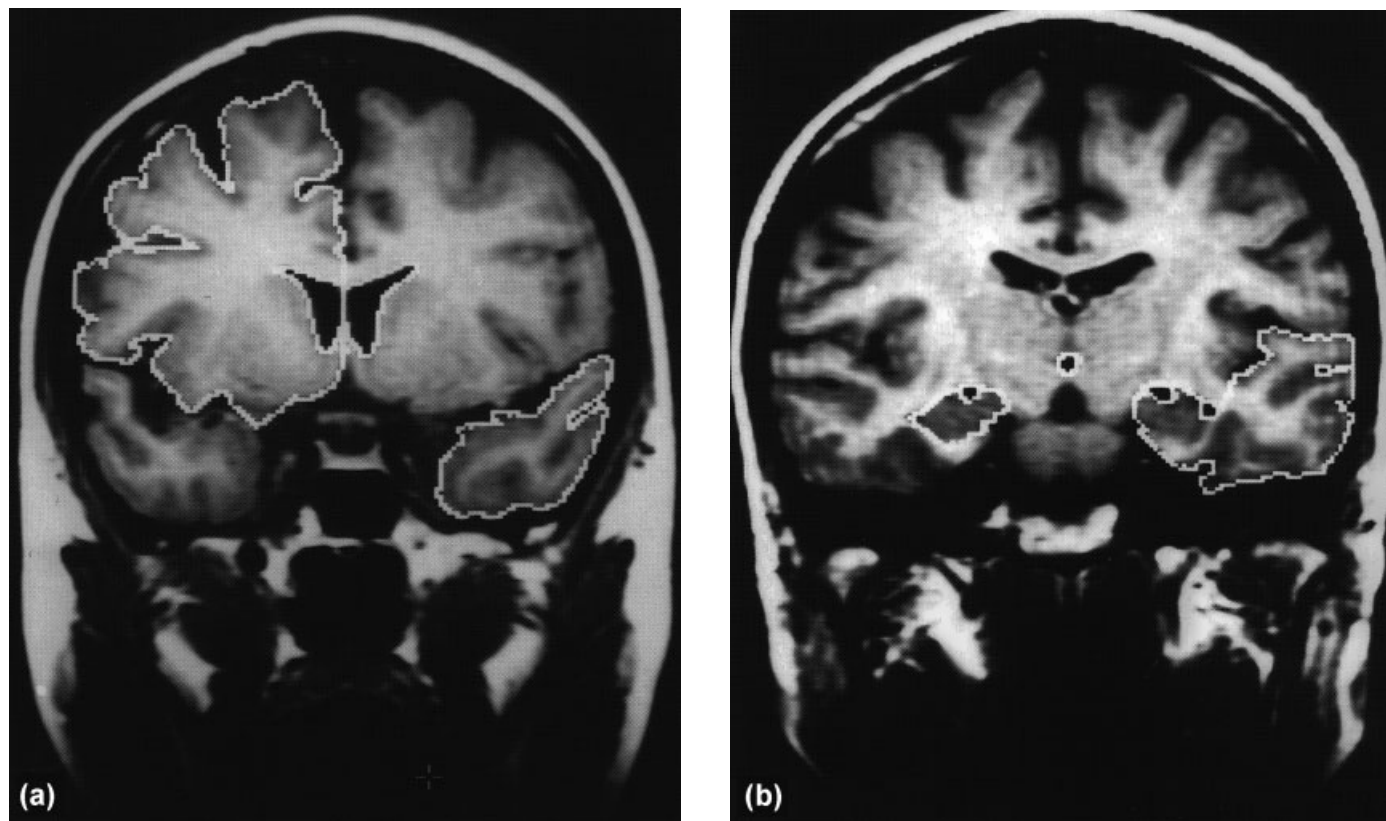


Figure 7.1 Typical coronal MRI images (TR = 500 ms, TE = 25 ms) illustrating anatomic landmarks and computer-assisted measurement of regional brain areas (A) Coronal section at level of optic chiasm, illustrating measurement of the right frontal lobe, the lateral ventricles and the left temporal lobe. (B) Coronal section at the level of the interpeduncular cistern, illustrating measurement of the left temporal lobe, the third ventricle and the right amygdala-hippocampal complex

an experienced research team that is blind to the subject's age and group status (i.e. normal control vs. patient).

These studies are ongoing, and several important findings have emerged^{41,99–104}. First, our quantitative imaging and measurement techniques provide highly accurate and reliable assessments of brain size. For example, intraclass correlation coefficients for interrater reliability range from 0.88 to 0.99, depending upon the particular brain region under study. The intraclass correlation coefficients for intrarater reliability range from 0.93 to 0.99. Second, regional brain volume clearly changes with age, with a predilection for the frontal lobes. We have observed a rate of decline of about 0.23%/year for cerebral hemisphere volume. Yet the rate of decrease we have observed for the frontal lobes is twice as great (0.55%/year). Our data also indicate that age related changes in regional cerebral volume is greater for ventricular regions than for parenchymal regions (3%/year vs. 0.23–0.55%/year). Thus, ventricular enlargement may prove to be a more sensitive index of brain aging than cortical atrophy. Third, formal assessments of cortical atrophy and ventricular enlargement suggest that the statistically significant age-related changes in brain volume may not be clinically significant. That is to say, anything more than mild cortical atrophy or mild ventricular enlargement appears to be distinctly uncommon in a medically healthy sample of elderly community volunteers. Fourth, the incidence of subcortical hyperintensity increases with age, but again it is uncommon for such changes to be severe^{39,96}. Fifth, the relationship between aging and changes in brain size is not a

simple one, in that it may be affected by a number of covariates (e.g. sex, height, years of education) that can modify the main effects of aging³⁸. For instance, we recently found that age-related changes in brain size were significantly greater in men than women for peripheral (sulcal) CSF volume, the lateral (Sylvian) fissure CSF volume, and the parieto-occipital region area¹⁰³. In addition we have observed significant effects of years of formal education on peripheral CSF volume, a marker of cortical atrophy¹⁰⁴. In a sample of healthy elderly volunteers living independently in the community and Mini Mental State Examination (MMSE) scores of at least 24, each year of education was associated with an increase in peripheral CSF volume of 1.77 ml. These findings are consistent with the 'reserve hypothesis', which posits that education (or factors for which it is a surrogate) provides a protective buffer against the injurious effects of age-related brain changes.

SUMMARY

Advanced brain imaging techniques produce highly accurate anatomic information and provide exciting opportunities to examine *in vivo* the effects of aging on the human brain. Our review of the existing literature indicates that normal aging is associated with cortical and subcortical atrophy, enlargement of the lateral and third ventricles, and an increase in subcortical hyperintensities. The extent of these anatomic changes can now be

quantitated with computer-assisted techniques and predictions can be made regarding the amount of change in tissue structure with advancing age. An important goal of future research will be to relate these anatomic changes to functional, neuropsychological and brain metabolic changes that accompany normal aging and age-related diseases of the brain.

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Quantitative Structural Changes in the Ageing Brain

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Estimates of the total number of human neocortical cells in both hemispheres range from 10 to 100 billions (10^9), but only two major studies describe how to estimate total neuron number. Reduction in the number of cells per unit volume of human cerebral cortex with age has been found in seven distinct neocortical brain regions from newborn to 95 year-old individuals². Haug³ estimated neuron density in 10–20 rows arranged perpendicular to the pial surface of the cortex in two to four areas per brain, multiplied by the assumed neocortical volume, and found an average of 15×10^9 neurons in the human neocortex. Haug found no decrease in total neuronal number with age and no difference in total numbers between males and females.

The three steps in estimating the total neuronal number in any defined brain region include: (a) delineation and estimation of the region's volume; (b) uniform sampling of the complex and irregularly shaped neocortex; and (c) estimation of the neuronal numerical density.

Based on unbiased principles, the Cavalieri method for the estimation of volume using systematic sampling and point-counting can be applied to any organ that can be cut into slices—physically or by means of optical or other scanning devices—and is independent of size or shape of the organ^{4,5}. In a recent study by Regeur⁶, neocortical volume, cortical thickness and volume of archicortex, the ventricular system, the central grey matter and white matter were estimated using stereological methods on brains from 28 old females (mean age 81.8 years) with increasing degrees of senile dementia, and brains from 13 (mean age 82.7 years) non-demented control females. Brains from the demented patients [14 Alzheimer's disease (AD) cases and 14 non-AD cases] had a smaller cortex volume and neocortical thickness was significantly reduced in the demented patients, with the highest degree of reduction in the most demented patients, as were the volumes of archicortex. No statistically significant differences were found in the volumes of cortex, white matter, central grey structures, ventricular volume or archicortex between the AD demented cases compared with the non-AD demented cases. The ventricular volume increased with increasing degree of dementia, but the difference between the demented group and the control group did not reach statistical significance. Surface area did not change in the demented patients, and no significant reductions were found in the volumes of white matter or central grey matter structures in the demented patients compared with controls.

In 1984 the dissector method was described, in which three-dimensional particles (e.g. neurons) are sampled with a constant

probability without regard to size, shape and orientation of the particles, provided that two requirements are fulfilled: (a) the complete set of particle profiles hit by the dissector's planar transects should be identifiable; (2) the dissector positions are uniformly random in the complete reference region⁷. The method relies on the fact that one can, without any other assumptions at all, count a cell nucleus or any defined particle within a defined reference volume if it is present in a relatively thin section but not in the previous member of a pair of adjacent sections. Particles of arbitrary size and shape can only be sampled with equal probability using a three-dimensional probe, such as the dissector. Systematically sampling a subset of dissectors from the wide range possible in a serially sectioned brain region or central nucleus, taking each with a constant but arbitrary probability, is sufficient for the estimation of the total particle number or, as in this instance, nerve cells (for practical details and analyses of sampling design, *see* refs 5, 8–11). Williams and Rakic^{12,13} have applied the same technique using the three-dimensional counting frame^{14,15} to obtain estimates of neuronal numerical densities.

A method for uniform sampling in human neocortex combined with a modification of the dissector principle, the optical dissector, has been introduced¹⁶. With the optical dissector the procedure is performed in thick rather than thin sections, which makes it many times more efficient. Using this method, 94 normal brains were studied, 32 females and 62 males in age-groups 20–90 years¹⁷. An unbiased estimate of the total number of neurons in the neocortex was obtained simply by multiplying the Cavalieri estimate of the neocortical reference volume by the numerical density obtained with the optical dissector. The number of neocortical neurons in females was 19.3 billion, and in males 22.8 billion, a difference of 16%. The difference in the total number of neocortical nerve cells over the observed range of 70 years was 9.5%, providing an average "loss" of neurons of about 85 000/day. This possible age effect was the same for both sexes. The total number of neocortical neurons in the material varied more than a factor of 2 with a range of 118% (14.7–32.0 billion neurons) (*see* Figure 8.1). The natural variability in neuron number of 19% ($CV = SD/mean = 0.19$) in normal Danes thus represents a variance of more than eight times the variance of body height [$CV(\text{height}) = 0.065$]: $(0.19/0.065)^2 \approx 8$.

On average, there were 186 million more neurons in the left hemisphere than in the right and this difference was the same in the two sexes. Sex differences were found in the total volume, total surface and thickness of the neocortex, white matter volume, central grey structure and brain weight. With advanced age,

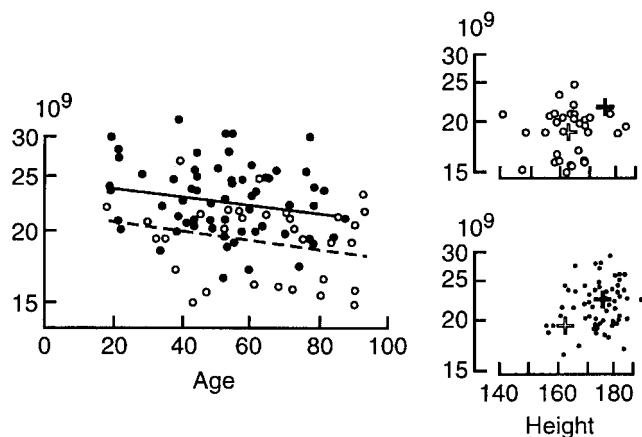


Figure 8.1 The number of neocortical neurons in normal Danes as a function of sex (●, males, ○, females) and age (left). The two orthogonal regression lines are indicated. To the right is illustrated the (absence of a) relation between the total number of neurons and the body height (cm) for each sex separately. The bivariate means for both sexes are shown in each diagram. All axes except the age axis are logarithmic

reductions occurred in neocortical volume, surface area, white matter, archicortex volume and brain weight, concomitant with a large increase in the ventricular system, while no change was found in grey matter volume and neocortical thickness. After sex and age were accounted for, neocortical neuron number was a dominating factor in determining the size of other brain structures. Neuronal density was not a function of sex or age.

A major problem in the interpretation of these data is evidently that one must take secular changes into account. Body height in Danish males has increased by approximately 9–10 cm from 1920 to 1980. Precisely *how* to correct for such changes is not known.

In conclusion, age may account for changes in both neocortical neuron number and neocortical volume without any effect on neuronal density. The reduction in cortical volume is seen without concomitant reduction in neocortical thickness, but only with consequence for surface area, a condition rather different from, for example, AD and AIDS, where the equally large atrophy only affects the neocortical thickness¹⁷. The largest changes in brain volumes are found in the brain white matters, with a reduction of $\approx 30\%$. In a recent paper by Tang *et al.*¹⁸, the total length of the myelinated fibres in five elderly women of 86 000 km was statistically significantly decreased by 27%, compared with 118 000 km in five younger females. As expected, the ventricular volumes increased by 50%, which could at least in part be a white matter volume reduction.

More research is needed to give us knowledge of possible changes during development, ageing and disease. Serious development defects and diseases, such as mental retardation, AD, schizophrenia and AIDS, have all been shown to involve

permanent structural brain changes. More information will help us to be able to separate the changes in normal ageing from those of neurodegenerative disorders and thereby understand the age conditioned functional deficits.

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Potential Regeneration of the Ageing Brain

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The brain does not spontaneously regenerate. As long ago as 1928, Cajal recorded that “once development was ended, the founts of growth and regeneration of axons and dendrites dried up irrevocably. In adult centres . . . everything may die, nothing may be regenerated”¹. Although it remains the case that the damaged mammalian central nervous system (CNS) does not generally generate new nerve cells in response to disease or injury, the intervening 70 years have identified a considerable plasticity of axons to remodel nerve connections and a limited degree of neurogenesis, which opens new opportunities for promoting

regeneration and repair in the damaged, diseased or ageing nervous system (see Figure 1).

COLLATERAL SPROUTING

The first clear evidence that Cajal’s dictum was overly pessimistic in relation to the mammalian CNS came from the demonstration that if a septal cell loses some of its normal axonal inputs, then other afferent axons can sprout into the vacated spaces to form

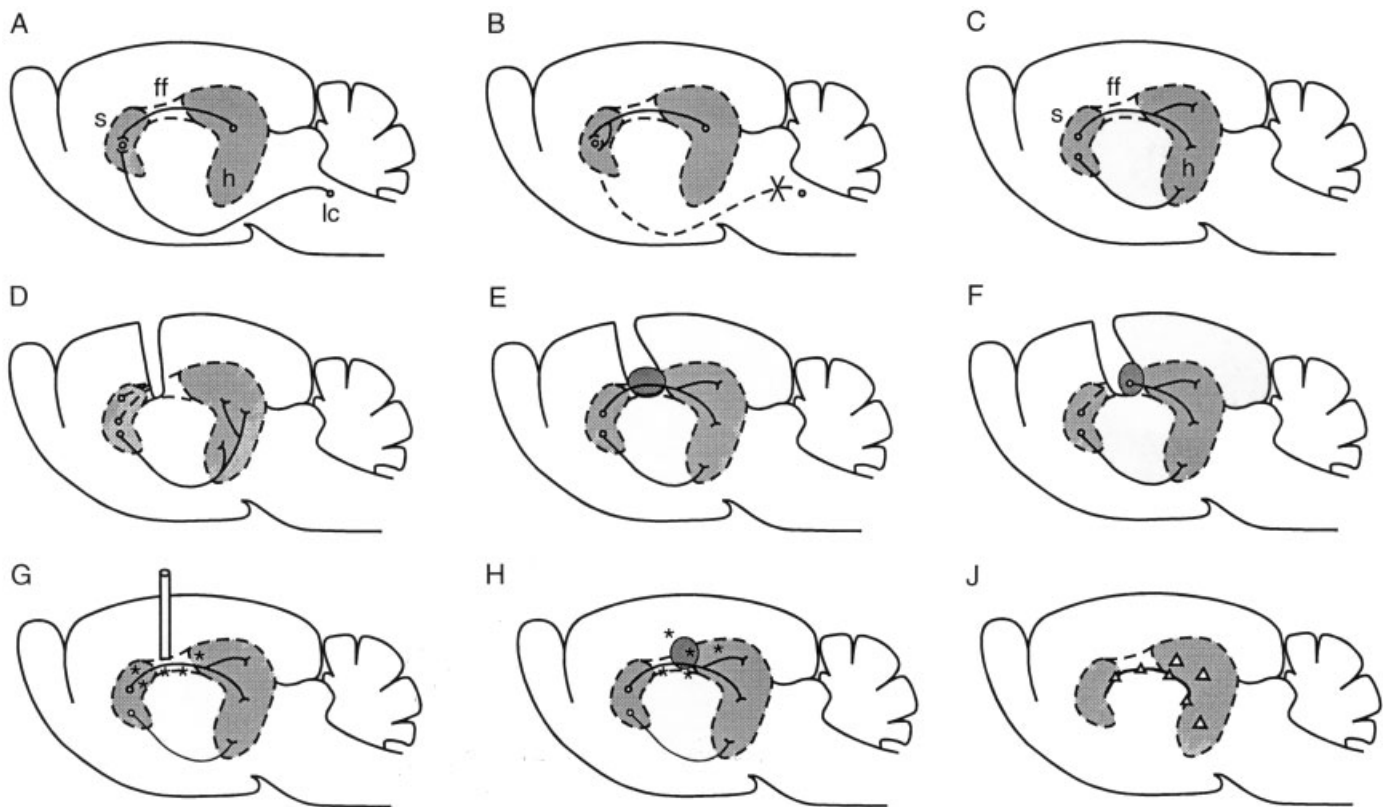


Figure 1. Schematic illustration of examples of regeneration and repair in the septo-hippocampal system of rats. (A) Normal synaptic inputs onto septal cells arising in the hippocampus and brainstem locus coeruleus. (B) Collateral sprouting of afferents from hippocampus following removal of afferents from the brainstem by a lesion of the medial forebrain bundle. (C) Normal septal projections into the hippocampus via dorsal and ventral pathways. (D) Compensatory collateral sprouting of ventral afferents following lesion of afferents to the dorsal hippocampus by transection of the fimbria-fornix bundle. (E) Implantation of hippocampal or glial grafts provides a substrate for regenerative growth of host septal axons back to the hippocampus. (F) Implantation of septal grafts provides a cholinergic reinnervation of the deafferented hippocampus. (G) Chronic injection of NGF molecules (triangles) diffusing via the lateral ventricles provides trophic support of cholinergic septo-hippocampal neurones in aged animals. (H) Implantation of cells engineered to secrete NGF provides similar neuroprotection in aged animals. (J) Neuronal stem cells in subventricular zone have the capacity to divide and differentiate in the adult brain. Whether they can also migrate to repopulate areas of cell loss in the aged brain remains speculative. ff, fimbria-fornix; h, hippocampus; lc, locus coeruleus; s, septum; small circles, neurones; small triangles, neuronal stem cells; asterisks, growth factor molecules

new synaptic connections with the target cell (Figure 1A,B)². If the new inputs are from a different source, they are by and large not functional. However, collateral sprouting of spared fibres of the same systems can sustain functional recovery in a number of model circuits of the brain (Figure 1C,D)^{3,4}.

REGENERATIVE SPROUTING

A major problem for extensive axonal reorganization in the adult brain is that although axons can undergo a degree of local sprouting, they do not typically retain the developmental capacity for long-distance growth through the CNS to distant targets⁵. Long-distance growth of axons can nevertheless be promoted by providing an alternative substrate for growth, such as Schwann cells from the PNS, which can be used to bridge a gap caused by a lesion cutting a pathway or be implanted as a track along which new axons can grow (Figure 1E)⁶.

NEURAL TRANSPLANTATION

When essential populations of neurons are lost, they may be replaced by transplantation. The techniques are now well established for transplantation of embryonic cells derived from the CNS into the aged brain of experimental animals (Figure 1F) and such grafts have been demonstrated to survive, repopulate areas of denervation, replace deficient innervations and restore lost functions in a wide variety of model systems⁷. There is now compelling clinical evidence that such grafts can provide a substantial alleviation of symptoms in Parkinson's disease⁸, and clinical trials are now underway in Huntington's disease, spinal cord injury and stroke⁹⁻¹¹. It remains a matter of speculation whether neuronal transplantation can alleviate the more diffuse and widespread degeneration associated with ageing and the dementias.

TROPHIC SUPPORT

Neuronal connections are dependent upon trophic support from their targets, and neurodegenerative diseases of ageing may in part be attributable to a decline in growth factor support¹². Thus, an alternative approach to prevention of progressive neurodegeneration, and to induction and guidance of regenerative axon growth, is to apply or replace identified trophic factors explicitly. For example, central cholinergic neurones, which decline in ageing, are dependent upon nerve growth factor (NGF) for trophic support. Prolonged injections of NGF into the ventricles of ageing rats can inhibit the progressive atrophy of septal cholinergic neurones and block the functional decline in the animals' learning abilities (Figure 1G)¹³. A similar strategy involving chronic central infusions of NGF has been attempted in a pilot experiment in Alzheimer's disease with only modest success¹⁴. A major issue to be resolved is how to deliver large trophic factor molecules—which do not cross the blood-brain barrier—to defined targets in the brain; the most powerful experimental techniques to date involve implantation of cells that are engineered to secrete the particular trophic factor molecule (Figure 1H)¹⁵.

ADULT NEUROGENESIS

It was believed until very recently that all neurones of the mature nervous system are born in early development, so that once lost they are not replaced. It is now clear that there exists in the adult

human brain a small population of resting "neuronal stem cells" with the capacity to both undergo further cell division and differentiate into both neurones and glia (Figure 1J)¹⁶. This opens the hope that such cells may be recruited for repair, either by isolation, expansion and differentiation into defined neuronal cell types *in vitro* and reimplantation, or by finding the means to induce their spontaneous division, migration to areas of cell loss and local differentiation into appropriate neuronal phenotype to replace lost target cells *in vivo*. However, it must be recognized that substantial technical and theoretical problems remain in translating such procedures into applications for repair in human ageing.

Although the repair of neurodegenerative damage associated with ageing remains experimental, rapid advances are being made in the techniques for inhibiting degeneration, promoting regenerative growth and replacing lost populations of cells. The pessimism that has for long surrounded the poor prognosis of brain damage in ageing and disease is being transformed to an optimism that these novel experimental approaches may find direct clinical application in the neurodegenerative diseases of ageing. Nevertheless, formidable technical problems still need to be overcome to transform theoretical prospects into practical therapies.

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Hippocampal Changes and Memory Impairment in Normal People

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INTRODUCTION

The nature of age-related cognitive deficits and their relationship to what is "normal", such as Alzheimer's disease (AD), remains a controversial area. It has long been recognized that decline in many aspects of cognitive performance occurs in the majority of individuals as they age. Most interest has focused on memory and various labels have been used, including "benign senescent forgetfulness (BSF)", "age-associated memory impairment (AAMI)", "aging-associated cognitive decline (AACD)"¹ and, most recently, "mild cognitive impairment (MCI)"². MCI describes the condition whereby a subjective memory complaint is accompanied by objective evidence of deficits, usually 1.5 standard deviations below age-corrected norms for a standardized test. Such individuals are known to "convert" to clear AD at a rate of about 15% per year³, in contrast to previous categories, such as BSF, when progression over time does not occur⁴. However, potential entities such as BSF, AAMI, AACD and MCI are still the subject of criticism, since it is unknown whether they represent a true disorder or a spectrum of the normal population mixed with those with early, and as yet undiagnosed, AD^{5,6}.

ROLE OF THE HIPPOCAMPUS

The hippocampus has long been known to be central to the human ability to learn new information, particularly for so-called episodic memory, i.e. memory for discrete events, and declarative memory, which is memory requiring conscious retrieval⁷. This contrasts with the role of the hippocampus in animals, which is known to be primarily involved in the formation of spatial memories. Recent evidence suggests that in humans the hippocampus does seem important in the acquisition of new spatial memories, as well as episodic memory, but is not the actual site of such spatial maps once they have been formed⁸. Differential functional organization of the hippocampus in time and space has been demonstrated⁷, giving support to the view that the main role of the hippocampus in memory formation is in forming new memories by capturing the event itself, coding it in time and space and binding it together for subsequent processing. Other brain areas (particularly the parahippocampal cortex and the frontal cortex) are also important in forming new memories, especially for encoding⁹. In contrast, other aspects of human memory (e.g. semantic memory) appears not to be dependent on the hippocampus¹⁰ although areas such as the anterior temporal pole and frontal lobe may be important¹¹.

RELATIONSHIP BETWEEN MEMORY IMPAIRMENT AND HIPPOCAMPAL CHANGES

While this has been reasonably well established for AD, the same is not yet true for using the memory impairment associated with ageing and conditions such as AACD, AAMI and MCI. An age-related decline in temporal lobe and hippocampal volume is found in most, although not all, studies¹²⁻¹⁴. Reductions in medial temporal lobe and hippocampal volumes and reduced perfusion

and functional imaging in normal individuals with memory impairment have been described by some groups^{15,16} but not others¹⁷, while the excellent spatial memory of taxi drivers has been related to enlargement of the posterior hippocampus in one study¹⁸. Further research is clearly needed. Similarly, some find a strong correlation between the degree of memory impairment in normal elderly people and volumetric change on magnetic resonance imaging, while others do not^{14,16,17}. More consistent is the finding that amongst those who already have memory impairment, reduced temporal and hippocampal volumes do predict those who will decline to develop dementia, as opposed to those who will not¹⁹⁻²². However, whether this is simply a reflection of the "contamination" of normal memory-impaired individuals and those with early AD remains unresolved.

Neuropathological data are limited, although MCI individuals may have Alzheimer-type pathology intermediate between those with AD and normals²³. Another candidate for the cause of any hippocampal damage during ageing would be excessive cortisol excretion associated with ageing, which is known to be toxic to the hippocampus in animals and possibly in humans²⁴⁻²⁶. These remain important avenues for future research. However, other possibilities exist for the neurobiological basis of memory and other cognitive impairments in healthy individuals. Generalized brain atrophy is known to occur with ageing^{27,28}, whilst an increase in white matter burden, presumably reflecting vascular pathology, occurs and has been linked with cognitive impairment in some studies²⁹.

CONCLUSION

Despite the hippocampus being central to new learning in humans and the evidence that memory impairment and hippocampal changes occur with ageing, further evidence linking the two is still needed. Undoubtedly, new views on hippocampal function and topographical organization, the better definition of the role of other structures, such as the parahippocampal gyrus and frontal lobe, in different memory functions, combined with further longitudinal clinical, electrophysiological, imaging and pathological studies, will provide a clearer answer to this important question in the near future.

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Neuroendocrinology of Ageing

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INTRODUCTION

Neuroendocrinology is the study of interactions between the nervous and endocrine systems. Such interactions occur via complex mechanisms involving the cerebral cortex, limbic system, brain stem, hypothalamus, pituitary and periphery with regulatory feedback by many hormones to the pituitary, hypothalamus and probably higher centres such as the hippocampus. Observations of neuroendocrine dysfunction and disorders of behaviour have helped to improve our understanding of mental illness and will be discussed later in this chapter.

Despite a surge of research over the last 25 years, many gaps remain in our understanding of neuroendocrinology, especially that of central secretion and neural control of hormone release. The complex interrelationships between monoamines, neuropeptides and hormones have not been elucidated¹. Hormones such as insulin and cortisol, once thought to exist only peripherally, are now believed to have central functions as neurotransmitters or neuromodulators². Neuropeptides such as opioids and vasoactive intestinal peptide (VIP) are found centrally and peripherally and some, such as thyrotropin releasing hormone (TRH), can be secreted from the same nerve terminals as classical neurotransmitters, thus also acting as neuromodulators³. Some neurotransmitters, once thought to act solely via the hypothalamus, may directly affect the pituitary, bypassing the hypothalamus⁴. The hypothalamus remains the fulcrum of neuroendocrine activity. In response to primarily central stimuli, the hypothalamus acts either: (a) via release of neuropeptide releasing or inhibiting factors into the portal circulation to the anterior pituitary; or (b) via the direct release of neuropeptides to the posterior pituitary. Thus adrenocorticotrophic hormone (ACTH), thyroid stimulating hormone (TSH), growth hormone (GH), follicle stimulating hormone (FSH), luteinizing hormone (LH) and prolactin are released from the anterior pituitary and vasopressin (antidiuretic hormone (ADH)) and oxytocin from the posterior pituitary. There is an intricate multilevel system of feedback mechanisms between adjacent and even distant steps in the process of hormonal secretion. Description of peripheral endocrine effects are beyond the scope of this chapter.

Like other systems, the neuroendocrine system undergoes a degree of decline during senescence, although by varying degrees and different mechanisms. Ageing may affect the neural control of hormones; the endocrine cells themselves, their hormones, hormone receptors and post-receptor events in target cells⁵. Most commonly there is a reduction in receptor cell numbers but hormone-receptor coupling mechanisms have also been implicated^{6,7}. As some symptoms of hormonal disorder

mimic the symptoms of advancing age it is not surprising that the hypothalamus and endocrine system have been linked as causative agents to the ageing process. Such organ system-based theories of ageing assume the presence of an "organal pace-maker" for ageing which initiates the chain of events seen in senescence⁸. The failure of organ systems, with the loss of homeostasis, is important to many of these theories. Some ideas have focused on more peripheral endocrine glands, e.g. the hypothyroid hypotheses, but there has been a more recent shift to central structures, such as the hypothalamus⁹. Dilman and others have proposed that an elevated hypothalamic threshold to negative feedback is an important element in the "deviation of homeostasis" after the completion of growth, and that a relative hypersecretion of, among others, growth hormone renders the internal environment inconsistent with survival by altering glucose and fatty acid metabolism and resulting in vascular disease, diabetes mellitus and other forms of morbidity¹⁰. Work by Frolkis *et al*¹¹ in Kiev has shown that the hypothalamo-pituitary axis plays a role in the regulation of RNA synthesis and in the induction of some enzymes of carbohydrate and protein metabolism. They found that these functions deteriorated with age and their work suggests that homeostatic and repair mechanisms may be influenced, which gives some support for an aetiological role for the neuroendocrine system in ageing. Possibly a more integrated theory involves a more circular mechanism. Genetic programming combines with the internal and external environment to result in impairment of all cells with ageing, including cells of the endocrine system. Impairment of the endocrine system alters homeostatic function and thus accelerates impairment of the remainder of the organism⁴. Given the complexity of the interrelationships within the internal environment, it is unlikely that the debate generated by these theories will be resolved in the near future.

Much of the initial research on ageing and endocrine function was with animal studies which, although helpful, may not be entirely applicable to humans, as there are important differences^{12,13}. However, techniques such as immunoassay have enabled us to measure hormone levels directly in humans, but even with this technique a high level of quality control is essential, as is the reliability of measurement. In human studies, too, we must remember that most samples are cross-sectional and we are therefore measuring age *differences* rather than age *changes*; longitudinal studies are therefore desirable. Nevertheless, in spite of these difficulties, greater sophistication and reliability in our measurement techniques, as well as the ability to synthesize hypothalamic releasing factors, have enhanced our understanding of neuroendocrinology and the effect of ageing.

AGEING AND NEURAL CONTROL OF ENDOCRINE FUNCTION

As previously mentioned, the neural control of hypophyseal secretion is complex and involves both classical neurotransmitters and neuropeptides, with good evidence that different neurotransmitter systems interact to modulate neurosecretion. Noradrenaline, dopamine, 5-hydroxytryptamine (5-HT), acetylcholine, γ -aminobutyric acid (GABA) and opioid peptides may all directly affect hypophyseal secretion mechanisms, whilst cholecystokinin, vasopressin and other neuropeptides have facilitatory or inhibitory roles¹⁴.

With ageing the secretion of neurotransmitters and receptor binding and *sensitivity* are altered¹⁵. Animal studies have suggested that the facilitatory effects of Ca^{2+} in neurotransmission may be reduced¹⁶. The overall brain content of noradrenaline is reduced, with many hypothalamic noradrenergic nuclei showing reduced noradrenaline concentrations with age, and adrenergic receptor sensitivity also appears to decline¹⁷. Dopamine receptor numbers are reduced, especially in the nigrostriatum. Acetylcholine metabolism also shows reduction¹⁸ and, although reports of reductions of brain 5-HT concentrations have not been fully substantiated, there does appear to be a reduced amplitude of circadian serotonin rhythm and a gradual decrease of 5-HT₂ receptors with age in the cerebral cortex¹⁹. It is likely that changes occur in other neurotransmitter systems with ageing, further altering the balance of neuronal control of endocrine function. In addition, neuronal sensitivity to feedback loop mechanisms may be altered, as reductions in the concentration of rat hippocampal cortisol receptors have been reported²⁰. Exactly how the effects of ageing upon neural mechanisms alter the functioning of subservient endocrine axes is not clear, but age-related changes in each neuroendocrine axis will be described below, with reference made to neural control mechanisms where appropriate.

AGEING AND HYPOTHALAMIC–PITUITARY PERIPHERAL ENDOCRINE AXIS

Morphologically age-related changes of the hypothalamus and pituitary are relatively minor. Calcification of the sella turcica may occur and animal studies have suggested an increase in intracellular lipofuscin, with a reduction in the volume of neuronal end swelling as well as in the number of hypothalamic neurosecretory granules²¹. Fibrosis with loss of basophil cells—hypophysitis navicularis—may occur in the anterior pituitary. Functionally, changes may be more pronounced and for the sake of clarity we will deal with each hormonal axis separately, as age-related change is differential.

HYPOTHALAMO–SOMATOTROPH–SOMATOMEDIN AXIS

The secretion of growth hormone (GH) declines in ageing humans²² with dampening of pulsatile GH release and loss of sleep- and exercise-induced swings. In addition, provocation tests suggest reduced GH reserve. The age-related changes, less pronounced in females, are due at least in part to reduced pituitary sensitivity to hypothalamic growth hormone releasing hormone (GHRH) and reduction of somatotroph numbers. However, reduction in sensitivity of α_2 receptors is also implicated, as evidenced by a blunted response to clonidine, an α_2 agonist²³. Peripheral α_2 receptor numbers have been found to be diminished. Altered catechol aminergic activity has been implicated in rats as administration of L-dopa reversed both mean and pulsatile GH levels²⁴. Somatomedin levels also fall, and it has been postulated

that reductions in GH and somatomedin may play a role in osteoporosis and other degenerative diseases in the elderly^{25,26}.

HYPOTHALAMO–HYPOPHYSEO–THYROID (HPT) AXIS

The HPT axis also undergoes changes coincident with ageing with a balanced decline in thyroid hormone production and degradation²⁷. In the elderly male, the overall secretion of thyroid stimulating hormone (TSH) is reduced and the response to thyrotropin releasing hormone (TRH) is blunted. This reduction is not marked and in females TSH levels may even increase with age²⁵. Pulsatility of TSH is maintained and, although results are inconsistent more recent findings have suggested that diurnal variation is also preserved²⁹. Healthy ambulatory ageing subjects show only a slight decrement in serum thyroxin (T4) levels. Thyroid binding globulin levels are unchanged but serum triiodothyronine (T3) concentration may be reduced by 10–20%, partly due to the age-related reduction in conversion of T4 to T3³⁰.

HYPOTHALAMIC–PITUITARY–GONODAL (HPG) AXIS

The HPG axis also undergoes gradual decline with age³¹. In men, primary testicular failure, possibly a result of reduced testicular perfusion, leads to a gradual reduction in testosterone levels, which becomes more marked after 60 years. Maximally stimulated testosterone, free testosterone and tissue testosterone are also reduced. Despite widespread variation, it is likely that such change is purely related rather than due to sedentariness or minor illness³². Reduced testosterone levels and elevated oestradiol (as a result of testosterone degradation in the skin and liver) may account for feminizing features, such as gynaecomastia, which may be seen in old men. Reduction of the feedback testosterone results in elevated levels of luteinizing hormone (LH) and follicle stimulating hormone (FSH) with age. However, sensitivity to negative feedback is increased at the hypothalamic level. The amplitude and frequency of pulsatile gonadotropin releasing hormone (GnRH) and luteinizing hormone releasing hormone (LHRH) are reduced, possibly as a result of neurotransmitter and neuromodulator changes, and pituitary sensitivity to releasing factors also appears to diminish. In ageing females, primary ovarian dysfunction occurs with shortening of the cycle length, depleted oestrogen and elevated FSH³³. Oestrogen levels fall to around 20% of the premenopausal level and then remain relatively stable, with the primary source of oestrogens being from androgen conversion in adipose tissue as ovarian androgens continue to be secreted³⁴. After the menopause, serum FSH and LH levels greatly increase to peak between 51 and 60 years and gradually decline thereafter. Animal studies have suggested increased pituitary sensitivity to LHRH but in humans it is not known whether the increased bioactivity of LH and FSH is as a result of greater pituitary sensitivity or high LHRH levels³⁵.

In both males and females, the production of adrenal androgens is reduced with senescence. Although prolactin levels are elevated in ageing rats and mice, it is unclear whether this is the case with humans, although recent evidence suggests a slight increase in elderly men and a slight fall in ageing women until the age of 80³⁶.

HYPOTHALAMIC–PITUITARY–ADRENAL (HPA) AXIS

The HPA axis is the best known and most studied of the neuroendocrine axes because of its intimate involvement in the

response to stress³⁷ (see below). Overall, findings suggest a mild reduction, if any, in basal functioning but with an altered regulatory capacity, an important factor in the stress response. Plasma total cortisol, plasma cortisol binding, plasma and urinary free cortisol and plasma ACTH are unchanged. Diurnal variation of cortisol secretion persists but may occur later in the day. Both ACTH and cortisol show decreased responsivity to provocative stimulation in older individuals and a degree of loss of the inhibitory feedback response³⁸. Several studies have shown attenuation of the cortisol response to ACTH, possibly as a result of altered functioning of ACTH-stimulated cAMP glucocorticoid receptors, whose numbers also diminish with age³⁹.

POSTERIOR PITUITARY

Morphologically, animal studies have revealed subcellular changes in the posterior pituitary. In humans, earlier evidence suggesting non-alteration of posterior pituitary hormones⁴⁰ must be tempered by the more recent evidence of a tendency to increased secretion of vasopressin in the elderly with elevated basal levels. Diminished secretion by ageing hypophyseal tissue is offset by elevation of vasopressin levels as a result of a reduction in renal tubule sensitivity to vasopressin⁴¹. Animal studies suggest that ageing may result in the loss of a particular group of hippocampal cells normally inhibitory to vasopressin secretion. Oxytocin levels also decrease with age.

AGEING AND STRESS

“Stress” may be defined in many ways but implies demands upon an organism threatening to overwhelm it and resulting in a physiological response. Human physiological responses to stress, either physical or psychological, are characteristic and involve initial adrenal medullary sympathetic activity which is later superseded by activity of the HPA axis during the “adaptation” phase. The HPA axis and its response to stress is outlined in Figure 9.1. Thus, in situations of stress, there is centrally stimulated hypersecretion of ACTH and cortisol, along with facilitatory vasopressin. The HPA axis also plays a major role in the homeostasis of stress and it has been postulated that it achieves this by blunting the organism’s persisting and potentially harmful physiological reaction to stress. The repeated aetiological linkage of stress to psychiatric illness has led over the last 25 years to much investigation of the HPA axis in stress and psychiatric illness, resulting in a mushrooming of the concept of psychoneuroendocrinology or behavioural endocrinology and the search for neuroendocrine markers for psychiatric illness. The concept of the neuroendocrine interface as “the window into the brain”, although not fulfilling initial expectations, has led to neuroendocrinological techniques becoming powerful research tools and valuable diagnostic aids in psychiatry⁴².

With ageing it appears that the physiological stress mechanism becomes compromised and there is also evidence that stressful stimuli can accelerate ageing⁴³. At the neurochemical level, rat experiments have suggested changes in monoamine metabolism and poor habituation to stress occurring with increasing age⁴⁴. Such a loss of habituation may be a result of an age-induced reduction of benzodiazepine binding sites⁴⁵ (thus increasing vulnerability to anxiety-induced mechanisms) or may be a consequence of a reduction of inhibitory hippocampal cortisol receptor numbers⁴⁶. Despite inconsistent findings there appears to be a diminished maximal response of the HPA axis to stressful stimuli, denoting a reduction in reserve capacity in ageing humans.

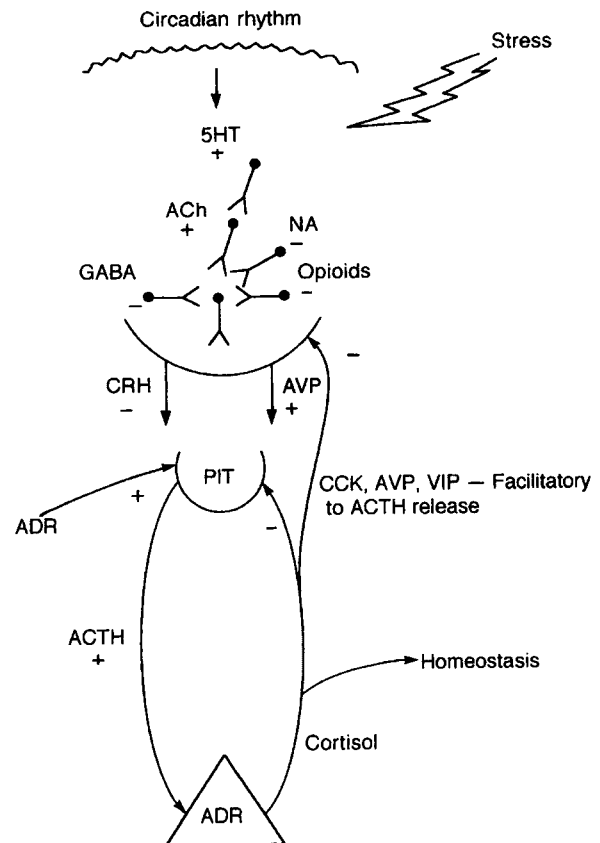


Figure 9.1 The HPA axis and its response to stress. 5-HT, 5-hydroxytryptamine; ACh, acetylcholine; NA, noradrenaline; GABA, γ -aminobutyric acid; CRH, corticotrophin releasing hormone; CCK, cholecystokinin; AVP, arginine vasopressin; VIP, vasoactive intestinal peptide; ACTH, adrenocorticotropic; ADR, adrenal gland

AGE AND PSYCHONEUROENDOCRINE MARKERS

As a result of research carried out over the last 25 years, it is now firmly established that mental illness is associated with a high incidence of endocrine abnormality in both young and old. The most commonly investigated endocrine axis in psychological disturbance is the HPA axis, which in depression characteristically shows hyperactivity with elevated circulating ACTH-cortisol and non-suppression by dexamethasone, a powerful synthetic steroid that, in normal individuals, inhibits the secretion of ACTH and hence cortisol. In younger individuals, dexamethasone non-suppression is not truly specific for depression and is common in other affective psychosis and acute schizophrenia, with a lower incidence in anxiety, panic disorder and anorexia nervosa⁴⁷. Dexamethasone suppression is also influenced by age (among other variables), with the healthy elderly showing a tendency towards non-suppression. However, non-suppression is more pronounced in the elderly depressed and those with Alzheimer’s disease. In the elderly, as in the young, the dexamethasone suppression test (DST) is still a useful aid when used with careful clinical assessment⁴⁸. It is helpful in differentiating depressive illness from minor psychiatric conditions and chronic schizophrenia, and reports have also suggested uses in identifying depressive pseudodementia and demented patients with depression⁴⁹. In depression, the DST is a good predictor of long-term outcome, with greater risk of relapse in non-suppressors⁵⁰. The

TSH response to TRH is also frequently blunted in depression but once again there is considerable overlap with other psychoses and there is an increased tendency to a blunted response in healthy senescent individuals⁵¹. Reports that blunted TRH responses are more pronounced in elderly depressives have not been entirely confirmed⁵². Growth hormone response to clonidine, an α_2 agonist, is blunted in depression, Alzheimer's disease and ageing. Recent data suggest that this blunting is again more pronounced in elderly mentally ill subjects than in the normal elderly⁵³.

CONCLUSION

Overall, although it is only in recent years that neuroendocrine function in the elderly has been studied in depth, it has been established that significant alterations in function occur with ageing in humans. Some of those changes may approximate endocrine abnormalities observed in younger individuals with mental illness, most notably depression. As yet our knowledge remains limited in the realm of behavioural neuroendocrinology, but such similarities in function of the aged and depressed have led to speculation of some common mechanism underlying age and mental illness. Hopefully research into such speculation will provide further enlightenment.

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Neurophysiology of Ageing as Reflected by Electroencephalogram (EEG) and Event-related Potentials (ERPs)

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INTRODUCTION

The electroencephalogram (EEG), which represents the electrical activity of the brain as recorded from electrodes placed at various positions on the scalp, is essential for the diagnosis of the epilepsies and the study of sleep disorders. It can also be a useful non-invasive aid for the detection and localization of structural brain abnormalities and the diagnosis of diffuse encephalopathies in patients with fluctuating levels of consciousness. Many of these conditions are common in the elderly population and it is therefore important to have a full understanding of the effects of normal ageing on the EEG, so that the significance of abnormal findings in elderly subjects can be more clearly interpreted.

ROUTINE EEG

Using standard procedures¹, electrodes, prepared with conductive jelly, are placed on the scalp and positioned in rows over frontal, temporal, parietal and occipital regions, according to the International 10-20 System². Most recording equipment permits an easy selection of different montages of electrode pairs and the filtered and amplified potential differences between pairs of electrodes are then recorded on paper trace (qEEG) and stored for quantitative analysis. Recordings are usually obtained from several scalp regions and under different physiological conditions. The resting EEG is recorded with the subject's eyes open and for a period with the eyes closed to test the responsiveness of background activity. Routine inspection of the EEG waveform takes account of changes in frequency, amplitude and response to activation procedures such as hyperventilation, which may be employed to accentuate certain EEG abnormalities. The presence or absence of paroxysmal activity will be noted. For the detection of brain abnormalities, changes in frequency are generally more reliable than changes in amplitude. By convention, four frequency bands are described: EEG waveforms in the range 8–13 Hz denote the alpha range, which is commonly observed in occipital brain regions of young persons during wakefulness when the eyes are closed (alpha is attenuated by visual attention). There is wide normal variation in the alpha rhythm, and slowing occurs with normal ageing, delirious states and metabolic disorders. If frequencies greater than 13 Hz are present, the EEG is said to show beta rhythm. Such fast activity may be found in normal people but is also increased by some drugs, including benzodia-

zepines and barbiturates. At the lower end of the spectrum, activity in the range 4–7 Hz is termed theta. Theta activity can be marked in young children, becoming less by the time of puberty. Frequencies below 4 Hz are grouped as delta activity. Theta and delta activity occur during sleep and are commonly found in neurological disorders whose slow wave abnormality may only be diffusely recorded, as, for example, in encephalopathies. Such slow wave bands also prevail when localized at electrodes overlaying space-occupying lesions.

THE ORIGIN OF THE EEG

The scalp-recorded rhythmic activity of the EEG is believed to be generated in the cerebral cortex, especially in large pyramidal neurones orientated vertically toward the surface of the scalp. However, rhythmic activity arising in subcortical regions, in particular in the thalamus, can be imposed on and modify the activity of these cortical cells via thalamo-cortical projections^{3,4}, so that the scalp-recorded EEG reflects changes in both cortical and sub-cortical structures. Much of the EEG variation usually detected can be attributed to hereditary factors. The early twin studies by Lennox *et al.*⁵, showing that brain electrical cerebral activity is strongly influenced by genetic factors, have been amply confirmed^{6,7}.

EEG CHANGES WITH AGEING

The most widely reported changes in the EEGs of elderly subjects are the slowing of alpha activity and the onset of focal theta and delta waves over the temporal regions^{8–12}. The changes are complex, as they affect the alpha, the slow wave and the fast frequency bands regarding both power and topographic distribution^{13–15}. Other reported changes corresponding to age include a much diminished slow wave response to hyperventilation and an increase in the occurrence of spike paroxysms in elderly subjects with no clinical evidence of a seizure disorder.

It is not clear, however, which if any of these changes are the result of a neuronal ageing process *per se*, rather than being manifestations of mild subclinical degenerative brain disease, including, for example, cerebrovascular disease, which is more common in the elderly. Quantitative EEG changes of power or complexity can be employed for highly successful statistical

discrimination between demented patients and elderly controls^{16,17}. Subtle cognitive impairment has been related to the presence of EEG abnormalities in some groups of otherwise healthy elderly subjects, supporting the view that many of the changes found in the EEGs of elderly subjects are due to specific subclinical pathologies¹⁸. No changes are observed in highly select groups of 'successfully aged' individuals^{14,19}.

NORMAL AGEING

In an early study on an elderly population, Silverman *et al.*²⁰ recorded EEGs of 90 healthy subjects aged over 60 and reported a diffuse slowing of the background rhythm in 26% and focal abnormalities in 43% of the subjects. These findings have been confirmed more recently^{10,21,22}. The slowing of the alpha rhythm with age was clearly demonstrated by Hughes and Cayaffa²³, who recorded the EEGs of 420 subjects aged 5–80 years. All subjects had been hospitalized and had undergone extensive neurological assessment in order to exclude the presence of brain pathology. In this group, the alpha peak frequency, which, up to the age of 60, had been between 10–11 Hz, fell to 9–10 Hz in subjects aged over 60. Some authors, however, claim a decreased slow and an increased fast activity with ageing²⁴. The annual changes in non-demented elderly are minimal compared to the alterations of alpha and theta power in demented individuals²⁵. There is no correlation between alpha and theta power and the degree of brain atrophy in the non-demented elderly²⁶.

Hubbard *et al.*²⁷ examined the EEGs of 10 centenarians aged 100–105, seven of whom were healthy, with no clinical evidence of degenerative brain disease. In this group, posterior dominant rhythms were in the lower part of the alpha range, and slow wave foci over temporal regions were common. These changes were similar to those found in subjects aged 80, and the study provided no evidence for a progressive decrease in alpha frequency or for an increase in focal temporal slow waves in subjects aged 80–100 years.

Changes in the EEG of elderly subjects, which have been attributed to early cerebrovascular insufficiency, include a diminished response of slow-wave activity to hyperventilation and the development of focal abnormalities, particularly over the anterior temporal regions. In a young person, hyperventilation for a period of 3 or 4 minutes usually produces a gradual increase in diffuse slow activity in the theta and delta range, which settles back to standard level within approximately 1 minute after cessation of over-breathing. This response is age-dependent and is most striking in children, who display delta activity at very high voltages. In contrast, old people show diminished or absent response to over-breathing, which may, in part, be due to diminished alteration in P_{CO_2} when hyperventilating⁴².

Bursts of rhythmic theta activity over the temporal regions frequently appear in late adulthood, and these are associated with cognitive and memory deterioration^{18,28}. In a recent study, Visser *et al.*²⁹ measured the EEG and performed computed tomography (CT) brain scans in a group of clinically healthy subjects aged 65–83 years. In this group of elderly subjects, those with focal EEG delta wave activity, recorded over the left anterior temporal region, performed poorly on neuropsychological tests of word fluency (thought to address temporal lobe function) and also had significant ventricular dilatation measured on the CT scan. It was concluded that such left-sided temporal slow-wave abnormalities found in the EEGs of some elderly subjects may be a valuable early indicator of temporal lobe pathology.

The probability that the EEGs of average adults do not change much throughout life and may, indeed, be relatively normal in otherwise healthy centenarians is thus raised²⁷. The slowing of EEG frequency with age could be explained by changes in

cerebral blood flow. Regional cerebral blood flow shows a strong inverse correlation with the appearance of EEG slow waves and is directly correlated with posterior alpha activity^{30,31}. A direct causal relationship between a reduced cerebral blood flow, an increase in slow waves and a reduced alpha frequency in the EEG of elderly subjects could, therefore, be postulated³².

EVENT-RELATED POTENTIALS (ERPs) AND AGEING

Electrical cerebral responses to discrete stimuli, such as visual, acoustic or contact stimuli, cannot, by and large, be detected in the scalp-recorded EEG. Electrical response to such events is small in comparison to their cerebral background activity, and averaging techniques are required for their visualization. Such techniques have proved of enormous value to neurologists and psychiatrists studying brainstem and higher cerebral function, by permitting the detection of tiny voltages generated in response to specific stimuli. To extract the time-locked activity generated by a given stimulus, a repeated series of stimuli is presented, and epochs of EEG, captured after each presentation, are summed and standardized/averaged. The random background EEG will tend to decrease in amplitude on summation, whereas the desired event-related potential will remain the same in size.

EXOGENEOUS EVENT-RELATED POTENTIALS

ERPs offer a means to assess peripheral nerve and brainstem function by using different sensory modalities. Early evoked potentials, generated within about 80 milliseconds (ms) after a stimulus, are described as exogenous because they seem to depend on the nature of the stimulus itself rather than any subjective response the subject may make to the stimulus. Auditory brainstem potentials, generated within the first 10 ms after a clicking sound, are evoked in a routine procedure to provide information about the functioning of auditory nerve and brainstem structures in the auditory pathway. Somatosensory event-related potentials (ERPs), evoked by electrical stimulation of, for example, the median nerve at the wrist, include the median nerve action potential, recorded at the brachial plexus, and activity generated in neurones of the spinal dorsal horn and dorsal column. Later peaks probably reflect activity in the medial lemniscus and primary somatosensory cortex. Early visual evoked responses to light flashes reflect activity in the visual path between the retina and the visual striate cortex.

From the second to the ninth decade of life, there is a linear increase in the latency of exogenous potentials³³. The latency of the median nerve compound action potential, recorded at the brachial plexus (the 'N10 waveform'), increases from an average of 10 ms in the second decade to approximately 12 ms in octogenarians. With few exceptions, a similar rise in latency with increasing age is found in all exogenous potentials of all three sensory modalities addressed and renders age corrections clinically important. The central conduction time of auditory evoked potentials (AEPs) increases by 1–4 ms/year, the latencies of visual evoked potentials (VEPs) by 2–4 ms/decade after age 40 years³⁴. Many age-related anatomical, physiological and biochemical changes may contribute to the slowing of nerve conduction implied by these latency delays.

ENDOGENEOUS EVENT-RELATED POTENTIALS

ERPs generated more than 80 ms after a stimulus may reflect the psychic condition of an individual. Such responses are termed

“endogenous” because their latency and amplitude are hardly influenced by the physical characteristics of the stimulus such as its intensity or frequency, but reflect how attentive the subject is. They also give an indication of the degree of complexity inherent in a cerebral cognitive or memory operation performed on exposure to a stimulus. The P300 response is one endogenous ERP that has been extensively studied, as it is thought to reflect the mental processes of selective attention, learning and memory. To generate a P300 response to an auditory stimulus, the subject is required to attend to a series of low-pitched (non-target) tones, randomly interspersed with high-pitched (target) tones. The recognition of these target tones generates a positive potential, which can be recorded widely over the scalp at approximately 300 ms after the auditory stimulus. The P300 response to the target stimuli is much more explicit, and it is generated only when the subject concentrates on the task; its amplitude is thought to reflect the level of attention and its latency the processing time involved in the recognition of a target tone.

EFFECT OF AGE ON LONG LATENCY EVENT-RELATED POTENTIALS

The latencies of all endogenous event-related potential components appear to increase with age from the second decade onward, a fact that led to extensive research activity in relation to the auditory P300 component³⁵⁻⁴¹. Some authors have reported a linear increase of P300 latency with age up to senescence with an increase of 1–2 ms/year. However, comparative studies, comprising large numbers of controls, have found an exponential ageing effect with a much higher rate of increase (of up to 4 ms/year) in P300 latency in subjects older than 60 years as compared to younger adults^{37,39,41}. P300 latency is also increased in the presence of a variety of brain pathologies, including the dementias of Alzheimer’s disease and cerebrovascular disease. In elderly subjects, it is difficult to separate the effect of ageing *per se* from the effects of subclinical degenerative or vascular changes on endogenous event-related potentials, and this is equally true for routine EEG. In a clinical environment, where P300 measurements may be useful, for example in dementia and schizophrenia, it is essential to carefully match individuals for age, particularly if elderly individuals are concerned.

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Pharmacokinetic and Pharmacodynamic Considerations in Old Age Psychopharmacology

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The elderly have the highest incidence of medical and psychiatric disorders. These conditions frequently occur simultaneously and are often chronic, lasting the lifetime of the individual. Consequently, the elderly require more medication than younger patients and even consumption of "over-the-counter" drugs is considerable among the aged¹. Because of this multiple drugs administration, they frequently experience adverse side effects. Psychotropic drugs have often been involved in such interactions, and cause twice the incidence of side effects in elderly patients as they do in younger patients¹. Drug interaction can produce a change in the pharmacological effect of a drug by altering activity at the site of action (a pharmacodynamic interaction), or by changing the plasma concentrations of a drug (a pharmacokinetic interaction), or both.

On the other hand, there has been a tendency of drug studies to focus on younger age groups and to exclude patients with comorbidity or polypharmacy². As a result, the generalization of current drug trials is a problem in old age psychiatry. Moreover, older adults respond less predictably than younger adults to most medication and this unpredictability is particularly evident among the frail elderly, who often suffer from central neurodegenerative disorders³.

For these reasons, physicians treating patients with multiple medications may be overly concerned about the potential risks of treatment and deny the patient the chance of recovery⁴. Physicians should, however, remember that response is generally good for elderly people who have specific psychiatric disorders, as is the case with major depressive episodes, and that the risks of leaving the client untreated could be greater than the potential risks of treatment.

Although treatment of the elderly may be complex, with some knowledge of both the pharmacodynamics and pharmacokinetics basics it is something that is both manageable and rewarding. In this chapter, the changes in the effect of psychotropic drugs with aging in general terms will be highlighted.

PHARMACOKINETICS

With aging, changes can occur in one or more of the different pharmacokinetic parameters; absorption, distribution, metabolism and excretion.

Absorption may be slower in the elderly or delayed in onset and this is due to several factors: reduction in gastric pH, diminution of the size of the intestinal absorption area, reduced mesenteric

blood flow (in general, the flow to all organs is diminished because of a decreased cardiac output). As a final consequence, the oral bioavailability of drugs may decrease in the elderly⁵.

Distribution is grossly influenced by the composition of the body; the volume of distribution may be increased in elderly patients due to a greater percentage of adipose tissue⁶. Most of the psychotropic drugs (and especially sedatives, such as benzodiazepines) are stored in fat tissues and this constitutes a contributory factor for the significant elimination half-life increase that is usually observed in the elderly. In fact, the elimination half-life is determined by clearance (which is the rate of drug removal per unit of plasma concentration) and by volume of distribution itself. The higher the volume of distribution and/or the smaller the clearance, the longer will be the elimination half-life. Moreover, both the lean body mass and the bodily water⁷ are decreased, so that ethanol (which is usually distributed across the bodily fluids) will show a higher concentration in the elderly than in younger groups⁸. Plasma protein binding is decreased in the elderly¹ because albumin concentration falls significantly. As a result, an increase in the free (non-bound) drug fraction in the plasma (the one that is able to pass the blood-brain barrier, but also the one that can determine the possible side effects) is observed. On the other hand, distribution of drugs into the brain may be decreased simply because cerebral blood flow (especially in arteriosclerotic patients) is usually diminished.

Metabolism. For psychotropics (lithium being the only notable exception), the most important metabolic processes are carried out in the liver (and elimination of metabolites will be made with faeces). As happens with most organs, the hepatic blood flow is diminished with age. The *metabolic transformation* processes are carried out by the microsomal enzymes in two different ways: hydroxylation and demethylation (processes that are significantly reduced with age) and conjugation with glucuronic acid (a process that is relatively unaffected in the elderly). This explains, for instance, the observation that benzodiazepines such as oxazepam and lorazepam (their major metabolic pathways involve glucuronidation) do not show clinically significant changes in pharmacokinetics with age⁹. On the other hand, chlordiazepoxide, diazepam and all the other pronordiazepam-like compounds (i.e. those that have *N*-desmethyldiazepam as the major metabolite) will show an increase of elimination half-life with age. In fact, these drugs are not metabolized by glucuronidation in the liver and, as a result, administration of these compounds in the elderly may cause a prolongation of action after a single dose and delayed accumulation on multiple dosing in the elderly¹⁰.

Metabolic interactions usually take place in the liver and are of special interest, as multiple drugs prescription is common in advanced age. The genetically polymorphic cytochrome P450 2D6 (CYP2D6) is responsible for the metabolism of several psychotropics. CYP2D6 activity does not change with age¹¹. CYP2D6 activity may be impaired by inhibitors such as paroxetine and fluoxetine¹², which can result in non-linear plasma drug concentration kinetics, as well as drug interaction when other drugs metabolized by CYP2D6 (such as desipramine, nortriptyline, neuroleptics, carbamazepine) are co-administered. Differently from paroxetine, fluoxetine and norfluoxetine (fluoxetine major metabolite), with sertraline the same pharmacokinetic parameters are found in both the young and the aged. Moreover, sertraline has much weaker inhibitory effects on CYP2D6¹².

Average dose adjustments for the aged can be derived from a simple equation and mean pharmacokinetic parameters from older and younger adults. However, individual dose adjustments (large variations in the decline of organ functions is possible) can be obtained from the drug clearance in a particular patient, where clearance/fractional bioavailability may be calculated from the area under the curve (AUC) of the drug in question⁵.

Excretion. Advanced age reduces renal function⁶, with important implications for lithium prophylaxis. The most important pharmacokinetic change in old age is a decrease in the *excretory capacity* of the kidney, so that the elderly should be considered as renally insufficient patients⁵. Since lithium is excreted in the urine, guidelines for lithium prescription recommend using a single bedtime dosing regimen¹³. Due to these considerations, an implementation of specialized clinics to manage and monitor elderly patients maintained on lithium has been proposed and discussed.

PHARMACODYNAMICS

Provided that pharmacokinetic guidelines for these adjustments are taken into consideration, the same plasma concentration is achieved in the elderly as in the young adults. However, we are frequently confronted with pharmacodynamic changes in old age that alter sensitivity to drugs, irrespective of changes of drug disposition⁵.

The central nervous system (CNS) is especially vulnerable. For example, aging can alter the sensitivity of the GABA carrier to some anesthetics (e.g. propofol and etomidate¹⁴). Moreover, in a well-known study¹⁵ the effects of a single 10mg dose of nitrazepam were compared with a placebo in healthy young and old people. Elderly people made significantly more mistakes in the psychomotor tests than the young, despite similar plasma concentration and elimination half-lives in both groups. The difference is probably explained by an increased sensitivity of the aging brain to the action of nitrazepam. It has been proposed¹⁶ that, with advancing age, and prematurely in Alzheimer's dementia (AD), the declining mitochondrial ATP synthesis increases GABA synthesis (a factor possibly responsible for forebrain dystrophic axonal varicosities, losses of transmitter vesicles and swollen mitochondria, markers currently regarded as earliest signs of aging and AD). Moreover, the particular vulnerability of the elderly to sedatives could be explained by aging-related changes in the expression of the gamma (2S) and gamma (2L) subunits in various brain regions, which suggest the existence of aging-related changes in the sub-unit composition in the GABA-A receptors, which in turn might lead to changes in receptor pharmacology¹⁷. Lastly, it seems that the activity of GABA-A transaminase (an enzyme that degrades GABA to succinic semialdehyde) is inhibited, which results in elevation of GABA content in the brain in some age-related neuropsychiatric disorders such as AD¹⁸.

The elderly are particularly sensitive to drug-induced parkinsonism¹⁹, which can reflect decreases with age in *dopamine* (DA) turnover and the suggested^{20,21} age-dependent deficit of the dopaminergic system, presumably related to a reduced number/activity of nigrostriatal and mesolimbic neurons. In humans, an age-related decline of binding of a ligand for dopamine transporters, specifically to the striatum, has been found, at the rate of 6.6% per decade²².

The response to agents with strong anticholinergic properties (tricyclic antidepressants; classical antipsychotics) increases in old age and may be accompanied by impairment of intellectual capabilities, agitation and, ultimately, delirium²³. However, non-demented elderly patients with psychiatric problems seem to tolerate psychotropic drugs (with respect to the impact on their cognitive competency) much better than patients with AD²⁴. Age-related reduced responsivity of the cholinergic system in the hippocampus has been well documented²⁵, but also disturbances in GABAergic/cholinergic interaction may play a key role in age-related cognitive dysfunction²⁶. During aging, higher affinity nicotine binding in the frontal cortex and the hippocampal formation decreases and these reductions may predispose the neo- and archicortex to the loss of nicotine *acetylcholine* receptor proteins observed in age-associated neurodegenerative conditions²⁷.

Studies in humans and primates suggest that the aged brain is prone to the degeneration of the locus coeruleus (LC)²⁸; the ascending dorsal noradrenergic bundle of the LC is involved in cognitive processes such as memory, learning and selective attention. Moreover, a profound *noradrenaline* depletion in the pre-frontal cortex (an area involved in certain cognitive functions, such as prevention of distractibility by irrelevant stimuli) has been described in the elderly²⁹. Together with the partial loss of CNS noradrenergic neurons, a compensatory activation of remaining CNS noradrenergic neurons has been described³⁰, which can explain the enhanced responsiveness (both in normal older subjects and with patients with AD) to noradrenergic agents such as yohimbine³¹.

The central serotonergic system is also adversely affected by aging, so that it has been proposed that possible reduction in humans of 5-HT_{2A} receptors and *serotonin* reuptake sites may contribute to ethanol consumption, depression and cognitive dysfunctions frequently seen in the elderly. These changes may alter the effectiveness of serotonergic drugs^{32,33}.

During the normal process of aging a number of changes in the glutamatergic system (involved in processes such as motor behavior, cognition and emotion), and especially a decrease in the density of *glutamate* NMDA receptors, have been described³⁴. Glutamate interacts with other neurotransmitters to conform the substrates of specific circuits of the brain that are relevant to aging. Impairment of intracellular energy metabolism associated with hyperactivation of glutamate receptors may contribute to the neuronal death seen in neurodegenerative disorders³⁵; the extent of glutamate neurotoxicity in the hippocampus is highly age-dependent, with mature animals' hippocampi more vulnerable to glutamate-induced cell death³⁶.

On the whole, aging is associated with changes in the regional brain chemistry and the brain multi-chemical networking profile (MCNP). In fact, there is an increase in overall chemical correlation in MCNP within and across all brain regions with increased age. This increased correlation may reflect an adaptive or compensatory response (possibly related to the elongation of the dendrites with aging) to the reduced levels of regional brain chemicals³⁷. Lastly, a diminished efficiency of the homeostatic mechanisms has been described in the aging brain, in part because of the reduced activity of various neurotransmitter systems. Counter-regulatory processes are therefore reduced and reactions to drugs may be increased²³.

PRACTICAL ISSUES

Bearing in mind the above-described pharmacokinetic and pharmacodynamic changes of psychotropic drugs in the elderly, a few principles could be recommended as prescribing guidelines³⁸: one should become familiar with a number of preparations and preferably administer them; use as few drugs as possible (including drugs for non-psychiatric conditions); 'keep it as simple as possible': give written instructions; avoid depot-forms, the treatment should be started in low dosages (1/5 to 1/4 of average adult dosage) and slowly increased (no sooner than every 5–7 days³⁹); the maintenance dosage is about 1/3–1/2⁴⁰ of average adult dosage ('start low and go slow'), but some elderly patients might need and can tolerate full doses⁸. The times required to reach steady-state therapeutic levels are longer.

Moreover, it ought to be emphasized that some psychotropics are more suitable for the elderly than others. For the treatment of affective disorders, *tricyclic antidepressants* are efficacious and inexpensive, but *SSRIs* and newer antidepressants are better tolerated and safer in overdose⁴¹. With respect to the putative diminished 5-HT responsivity in this population, the ability to identify SSRI non-responders via 5-HT challenge in combination with neuroimaging measures may have important clinical utility³³. Among the SSRIs, preference is possibly given to sertraline¹². The selective *MAO-A inhibitors* have not been extensively studied in the elderly, but they have definitely overcome the use of the classical MAO inhibitors. *Lithium* is still the mainstay for the treatment of bipolar disorders, but careful dosage and monitoring of plasma concentration are necessarily required. On the other hand, bipolar elderly patient responders to *valproate* ought to achieve higher serum concentrations of valproate itself⁴².

The age-related changes in the pharmacokinetics and pharmacodynamics of the *benzodiazepines* (still the most frequently prescribed drugs for anxiety in the elderly) recommend preferential use of those agents that are metabolized via conjugation (oxazepam); *risperidone* (which is better tolerated in the elderly) may be used as an alternative. However, together with the sedation increase, with a sedative/hypnotic prescription a cognitive function decrease is observed in the elderly, with consequent risks of falls and injury (especially if the diazepam-equivalent dosage is higher and if the patient is prescribed with more different drugs)⁴³. According to some suggestions⁴⁴, because of the high level of comorbidity between generalized anxiety disorder and major depression in late life and the observation that anxiety is usually secondary to depression, antidepressants constitute the primary pharmacological treatment for many older people. For the treatment of insomnia, both zopiclone and temazepam are to be considered as effective hypnotics, but the first shows a superiority on sleep architecture⁴⁵. New promising agents, such as *cholecystokinin-B receptor antagonists*, seem to be specific, in aged animals, for an improvement of sleep quality⁴⁶. For the treatment of psychotic syndromes, due to the elderly extreme sensitivity to parkinsonian side effects and to the anticholinergic properties of the *classical antipsychotics*, attention is given to the *newer antipsychotics*, but there is still a paucity of data. Clozapine may be a useful drug but adverse effects can occur⁴⁷.

Notwithstanding the aforementioned I am in agreement with Jovic³⁸, who stated that "psychotropic drugs cannot compensate for the lack of human contacts, devotion and intensive relationships, but complement them".

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Normal Ageing—A Problematical Concept

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INTRODUCTION

The word “normal” is used to refer to what is statistically normal, that is within the average range. The range can vary somewhat, say from the middle 50% of a normal distribution to perhaps the middle 80%, depending upon one’s purpose. The word is also used to refer to prototypical members of a category, members with characteristics that best exemplify the category as a whole. The word is used to refer to what is socially prescribed and expected, such as the usual forms of appearance and behaviour for a given occasion in a community. Another usage refers to a standard pattern or sequence of events that have a high probability of occurrence, such as impairment of vision and hearing in later life.

NORMAL AND PATHOLOGICAL AGEING

With regard to human ageing, the word “normal” is used in all the above senses, depending upon the context. In professional gerontology, however, the phrase “normal ageing” usually implies the existence of a contrasting condition or process, viz. “pathological ageing”. Difficulties arise because normal ageing and pathological ageing are conceptually interdependent. The main historical landmark in attempts to distinguish between them was the publication of Korenchevsky’s *Physiological and Pathological Ageing*¹. Korenchevsky drew attention to the fact that some physiological functions in some elderly human subjects were equal to or superior to those of chronologically younger subjects. Psychological research into sensorimotor and cognitive performance often reveals that some elderly subjects perform as well as or better than the average younger subject.

On the basis of evidence that some individuals show relatively little physiological impairment with age, at least until late life, Korenchevsky inferred the existence of primary (non-pathological) ageing. On the basis of evidence that other individuals show substantially greater than average impairment earlier in adult life, he inferred the existence of secondary (pathological) ageing.

These two inferences, however, are simply two versions of the same argument, namely, that ageing is characterized by wide differences between individuals. If we plot the distributions of scores on physiological or psychological functions for several age groups in a cross-sectional study, we often find considerable overlap between even widely spaced age groups. If the distributions of performance scores for the same respondents at different ages in a longitudinal study are compared, we usually find that individuals tend to retain their position (rank order) relative to other respondents. A minority, however, show decline relative to

their position at earlier ages. These are the people who appear to exhibit pathological ageing. Thus, individual differences in normal ageing tend to be maintained, even though there is a decline with age, on average, over the period studied. These differences are brought about by various causes, including genetic characteristics, life-history events, life styles and environmental conditions.

Even among the community-dwelling elderly, there are wide variations in physical and mental health and wide variations in such things as living conditions, social support, stress and coping strategies and health. In a multicultural society, the range of differences between individuals at later ages is likely to be very wide. The process of normal ageing is a social as well as a biological process. That is to say, society *prescribes* or *normalizes* various stages in the life cycle, so that there are typical ages for the completion of full-time education, marriage (or sexual partnership), parenthood, occupational status and retirement. Such arrangements may change from one generation to the next. This, together with secular changes in health, longevity, life styles and so on, make the concept of “normal ageing” a moving target. Flynn has reported substantial secular (cohort) effects on measures of intelligence². Consider how the contraceptive pill and hormone replacement therapy have changed the life styles of women. Consider also how drugs, AIDS, migration and economic factors may affect ageing in sections of the population. These are technical issues for demography and epidemiology.

It is possible to demonstrate general age trends and effects. For example, the sex difference in longevity is well established; there is a differential decline in fluid and crystallized psychological abilities; anatomical and physiological functions have their characteristic normal patterns of change with age. There are some similarities between the normal (common) effects of ageing and the effects of pathologies such as Alzheimer’s disease, as shown by neurological and psychological tests. These trends and effects are compatible with the view that ageing is the result of a multiplicity of causes. They are not proof that there are two sorts of ageing: pathological and normal (non-pathological)³. On the other hand, there is the question of whether senile dementia of the Alzheimer type or multi-infarct dementia are the end-results of a normal intrinsic ageing process that would affect anyone who lived long enough, or whether they are abnormal conditions induced by genetic faults, life-history factors or specific extrinsic causes, such as infection or exposure to noxious substances. Genetic mutations increase with advancing age, and may affect performance before the obvious signs and symptoms of disease.

The argument in favour of the notion that there are two sorts of ageing—normal and pathological—is supported by evidence that people suffering from identifiable pathologies, such as cancer, heart disease or diabetes, have reduced life expectations and are

functionally less competent in some respects (see van Boxtel *et al.*⁵). Moreover, some of these disorders are age-related; some, such as Simmonds' disease, mimic the normal (usual) effects of ageing. Individuals who survive to a late age do not have a history of such disorders. The difficulty with this argument is that it is circular: pathological conditions are conditions that increase the likelihood of functional impairment and death; conditions that increase the likelihood of functional impairment and death are pathological. If an adverse effect commonly associated with age is not attributable to pathology, then, by definition as it were, it is "normal". If the underlying cause is identified, it is then labelled "pathological". Diseases can be regarded as concepts rather than entities (unless a cause can be found), in which case the distinction between normal and pathological ageing is a matter of definition, not an empirical issue. The empirical issue is how to identify and deal with the many age-related causes of impairment, regardless of whether they affect many people or just a few.

In order to demonstrate the existence of pathological ageing (as distinct from pathologies that increase functional impairment and the probability of death), we would need to show stepwise discontinuities in age trends, or departures from the "normal" distribution of differences in performance. Stepwise discontinuities and bimodal distributions are not common in the sorts of samples recruited for cross-sectional or longitudinal research in ageing.

Defining pathology in terms of a marked deviation from normal function means that the cumulative adverse effects of ageing eventually become pathological relative to standards for younger people but not older people—hence the view that there are many normal old people but few healthy ones! Stoller reported a tendency for older people to interpret their symptoms in terms of pain, discomfort and interference with their activities, rather than in terms of a possible medical condition⁵.

Improvements in living conditions, diet, exercise and medical treatment have the effect of extending the average span of life, and so have the effect of redefining what we mean by normal and pathological ageing⁶. The distinction between the "young old", and the "old old" is now well established. Normal ageing can be taken to mean that set of intrinsic age-related effects that characterizes the adult life of people who occupy the middle ground of a distribution of age at death, or that characterizes and explains the average elderly person's functional competence. Pathological ageing can be taken to refer to the intrinsic age-related effects that characterize people who die relatively young, or who perform well below comparable people of the same age, as a consequence of these effects. The problem here is to demonstrate a causal connection between age-related ailments and the so-called "intrinsic" effects of ageing.

Normal or intrinsic ageing can be regarded as a species-specific process of degeneration, subject to a degree of variability depending upon initial genetic endowment and subsequent environmental conditions. Pathological ageing is any substantial deviation from the normal (standard or common) pattern of age-related changes.

THE CAUSES OF AGEING

When we examine the distribution of age at death for human populations, we find a relatively "normal" distribution on which is superimposed a "tail" representing infant mortality, accidents and premature deaths in early adult life. A normal distribution is typically the result of a multiplicity of independent contributory causes. The distributions of scores for physiological and psychological functions, obtained from reasonably large samples of the sort usually recruited in studies of ageing, tend to be relatively normal. So the assumption is that these effects, too, are the result of a multiplicity of independent causes. The aim of such

studies is to identify the factors that account for most of the observed variation^{7,8}. In some research studies, gender, education, health and intelligence account for a substantial part of the variation in psychological performance, leaving chronological age accounting for little.

The existence of trends and effects associated with chronological age does not imply a causal agent—"ageing". As we have seen, ageing is simply a convenient label for a variety of age-related primary and secondary causes of impairment. Some of these causes no doubt interact and produce many sorts of indirect and long-term effects. For example, injuries, stresses, learning experiences and many other eventualities can occur at different ages. They may have different consequences because other age-related changes have or have not taken place, and because of differences in people's biological, psychological and social characteristics. In any event, given the complexity of the processes involved in human ageing and the long periods of time over which the processes occur, a considerable amount of "turbulence" in the effects of ageing is to be expected. Even small differences in initial personal characteristics and circumstances could lead to wide differences, in physical and mental health and performance, between individuals at later ages.

On this view, older people who are physiologically and psychologically very competent are simply those who have a superior biological constitution, have suffered fewer or less serious adverse effects from the many age-related causes of impairment, and have been able to take advantage of circumstances that promote health and well-being. People who survive to very late life with high levels of physiological and psychological competence are sometimes referred to as a "biological elite". Such people occupy, say, the top 10% of a distribution of physiological or psychological performance. They are studied retrospectively in the hope of identifying some of the factors that contribute to longevity and functional competence in late life.

Older people at the other end of the distribution are those who have not had the same benefits, and are regarded as suffering from one or other of the disorders of late life. On this view, the elderly who occupy either the upper or the lower tail of a relatively smooth normal distribution are different only in degree, not in kind, from those that occupy the middle ground. Of course, future research may identify specific positive and negative factors that will help us to account further for the observed variations in longevity, health and performance.

BASIC RESEARCH AND INTERVENTION

Health, education and gender are important variables in basic research in ageing. Controlling for them often substantially reduces the effects of chronological age (the usual index of "ageing"). It is possible to contrast the age trends in physiological or psychological performance for subjects suffering from known pathologies with the age trends for subjects free from such pathologies. Rabbitt⁹ reports that when adults with diabetes are compared with subjects who do not have diabetes but are comparable in other respects, including intelligence, the diabetics perform less well on measures of information processing. Rabbitt also reports that moderate deafness can have a similar, indirect, deleterious effect on information processing. This supports the use of specific health measures as co-variables in research on ageing.

One of the difficulties in both cross-sectional and longitudinal studies of ageing is that samples of older respondents are biased in at least two ways. First, volunteer respondents tend to be physically, psychologically and socially advantaged, relative to non-volunteers of the same age. Sampling bias is a major obstacle to research in ageing. It restricts our ability to generalize our findings to the wider population. Second, older respondents, even

older volunteers, are more likely than younger respondents to have various physical and psychological impairments that adversely affect their performance: for example, lack of exercise may impair cardiac function; poorer vision or hearing may slow reaction times and increase errors; social isolation may increase anxiety or depression; disuse, as well as irreversible deterioration, may impair performance, as in driving or playing games; training and practice can help compensate for these losses. These and other sampling biases mean that cross-sectional or longitudinal comparisons are likely to be distorted. If we do not have data on relevant background variables, then we cannot take them into account when we try to interpret age-related effects on performance.

A distinction can be drawn between research samples that are actually representative of their age group, and samples of other sorts. People who survive to later ages differ in a number of ways—biologically, psychologically and socially—from non-survivors. Thus, a representative sample drawn from an older population is not directly comparable with a representative sample drawn from a younger population. Matching members of an older sample with members of a younger sample means using samples biased with respect to the parent population. In a longitudinal study, dropouts have the effect of changing the population represented by the remaining sample.

Research samples consist of two main sorts: (a) volunteers recruited by advertising, “snowballing” or other forms of gentle persuasion; and (b) subjects recruited in ways that make it difficult for them not to take part. In the latter sort of recruiting, the benefits of participation are held to be of direct personal benefit, or in the public interest, as with samples based on medical registers. Risch *et al.*¹⁰ report on the difficulties encountered in recruiting “normal” volunteers for psychiatric research. Systematic psychiatric screening, over the telephone, failed to exclude 25% of unsatisfactory volunteers, as judged by subsequent tests. However, in a smaller subsequent study, a warning that reimbursement would be withheld if their toxicology test proved positive was effective. In reply, Halbreich confirms the need for close examination of research volunteers, but raises questions about how to achieve this, and about the ethical issues involved¹¹. He also considers that screening normal volunteers should include a family history.

Most psychological studies of ageing appear to assume, rather than test, that their volunteers are “normal”, although there may be routine mention of the fact that the respondents are free of obvious physical or mental impairment. The usual assumption is that volunteer samples are biased towards better-than-average health, ability, education and social status. Todd *et al.*¹² describe ways of improving volunteer rates. Bromley (*op. cit.*) describes a number of methodological difficulties encountered in research into normal ageing, including the difficulties associated with sampling and psychological measurement.

The effects of age, cohort and time of measurement are difficult to disentangle. The method of age-matched controls is often used to compare normal with pathological ageing. The problem here, as in selecting a sample of normal respondents, is how to determine which variables need to be controlled. There are numerous biological, psychological, social and environmental factors known or thought to be involved in ageing, and doubtless many unknown factors. The onus is on the investigator to justify the inclusion or exclusion of particular controls, and on the critic to identify other possibilities. Research needs to move beyond superficial and crude controls for gender, socio-economic status and self-reported health. How this move is to be achieved is itself a research issue (*see e.g.* Fox *et al.*¹³). Social groups that have experienced specific long-term life styles, for example in relation to diet, exercise, exposure to toxic substances, can be compared with “normal” social groups in relation to ageing.

Interventions designed to improve the functional capacities of the elderly have had some success but leave some questions unanswered. A sense of control and efficacy appears to be an important and modifiable personality characteristic in later life¹⁴, but intervention raises ethical and management issues. Physical exercise is effective in improving physiological functions even late in life, but prospective studies are needed to exclude the effects of sample bias and lack of control subjects¹⁵. Rubin *et al.*¹⁶ describe a prospective study of the onset of dementia in apparently normal elderly volunteers.

Training and practice on sensorimotor and cognitive tasks improve the performance of elderly subjects, but improve the performance of younger adults too. The effects may not generalize much beyond the training task or persist long after training is discontinued. The main point, however, is that performance on the first occasion of an unfamiliar task (in a typical laboratory study) is not necessarily indicative of practised performance on a familiar task of the sort encountered in daily life^{17,18}. Recently, more interest has been shown in the role of crystallized abilities (acquired mental skills) in adult life¹⁹. Practice and experience help to explain the maintenance of high levels of performance in many normal old people.

The concept of normal ageing refers not only to humans but also to other animal species, such as flatworms, rats and monkeys. Plants, too, have their characteristic life cycles. Different strains within a species have their characteristic, normal, patterns of ageing. Selective breeding and treatments, such as dietary restriction and brain chemistry, are used to explore the causes of longevity and age-related pathologies.

It is impossible within the scope of a short article to do justice to the numerous and diverse publications on “normal ageing” that a computer-assisted literature search can identify. The concept of “normal ageing” can be used to refer to various phenomena. The key question is, “Normal in respect of what?”

CONCLUSION

Normal ageing can be defined as a cumulative process of adverse changes in physiological, psychological and social functions that, in a general way, characterize average members of successive older cohorts of adults. This process is at present irreversible and to some extent predictable, but it produces a wide range of differences between individuals in age of onset and rate of change. To a limited extent, people can retard and ameliorate these adverse changes.

Normal ageing is a “socially constructed” concept, referring to an accepted range of variation in the health, appearance and performance of adults at different ages. It is also a “scientifically constructed” concept, referring to research findings in gerontology and other disciplines. Gerontologists find the concept of pathological or abnormal ageing useful in identifying exceptions to and deviations from the normal pattern, but the distinction between normal and pathological ageing remains problematical.

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Cohort Studies

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The word “cohort” originally designated a Roman military unit but has now become a technical term in population studies as a collective noun for any peer group, band or sub-set of individuals under investigation. In current usage, a cohort is any group of individuals who are linked in some way. This link may be experience of common life events, such as particular pathologies, or life transitions, such as menopause, or experience of a particular historical event or socio-economic condition. In gerontology the most common defining factor is age group, and this should be assumed unless some other usage is specified. In gerontology, “cohort” has become a more acceptable term to specify chronological age than the often misused “generation”, which has an equally precise and different meaning in studies of kinship terminology.

Note that “age group” may be very loosely defined. For example cohort members may have all been born, or died, in the same week, month, year, decade or even century. The defining boundaries of cohorts cause methodological difficulties because the effects of age are confounded with those of birth cohort and period. That is, groups of people born at the same time (birth cohorts) are by definition all of the same age, and have all lived through the same historical period. Groups of the same age (age cohorts) are not necessarily born at the same time, and so may have experienced different historical periods and events such as wars, with attendant differences in social circumstances. Groups of different ages (different age cohorts) also have not shared particular historical events or periods and, to the extent that their experience of a historical event has overlapped, they have been affected by it at different ages.

Cohort analysis is the methodology of designing and analysing studies to make inferences about the behaviour or condition of a particular sub-group without the necessity for studying them again after one or more successive time periods. It is now the most

common methodology used to study changes in behaviour or attitudes, biological and cognitive effects of human ageing and social, political and cultural change. Cohort analyses in developmental and ageing studies are distinguished from longitudinal analyses, in which the same birth cohorts are re-examined at intervals over a period of time. A compromise is cross-sequential analyses, in which different age groups and birth cohorts are repeatedly re-examined and compared with themselves and with others at different measurement points.

Unfortunately, although cohort analyses always allow us to identify and study each of these effects, they do not provide any direct way of examining all of them, independently, in a single study. One reason for this is sampling variability. That is, any cohort samples we can obtain and compare are unlikely to be precisely comparable. A related difficulty, which usually guarantees that samples in gerontological comparisons will not be comparable, is sample attrition. As cohorts age, they lose members and so alter in terms of their credibility as representative samples of the populations from which they were initially selected. The most important limitation on cohort analyses is that there is no way to avoid confounding at least two of the three variables in which we are usually interested: age, cohort and period. This imposes very inconvenient restrictions on statistical analysis; for example, in a multivariate regression analysis, all three variables cannot simultaneously be entered as variables in a regression equation. Although many attempts have been made, and much has been written on the subject, there are still no statistical methods that can clearly separate the effects of birth cohort and period from the effects of ageing. Perhaps the most lucid and helpful discussions have been by Costa and McCrae¹ and Palmore². The current consensus is that decisions must rest on scientific judgement as to which two of the three possible effects are likely to be important, so that the third can be omitted. The

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Cohort Studies

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The word “cohort” originally designated a Roman military unit but has now become a technical term in population studies as a collective noun for any peer group, band or sub-set of individuals under investigation. In current usage, a cohort is any group of individuals who are linked in some way. This link may be experience of common life events, such as particular pathologies, or life transitions, such as menopause, or experience of a particular historical event or socio-economic condition. In gerontology the most common defining factor is age group, and this should be assumed unless some other usage is specified. In gerontology, “cohort” has become a more acceptable term to specify chronological age than the often misused “generation”, which has an equally precise and different meaning in studies of kinship terminology.

Note that “age group” may be very loosely defined. For example cohort members may have all been born, or died, in the same week, month, year, decade or even century. The defining boundaries of cohorts cause methodological difficulties because the effects of age are confounded with those of birth cohort and period. That is, groups of people born at the same time (birth cohorts) are by definition all of the same age, and have all lived through the same historical period. Groups of the same age (age cohorts) are not necessarily born at the same time, and so may have experienced different historical periods and events such as wars, with attendant differences in social circumstances. Groups of different ages (different age cohorts) also have not shared particular historical events or periods and, to the extent that their experience of a historical event has overlapped, they have been affected by it at different ages.

Cohort analysis is the methodology of designing and analysing studies to make inferences about the behaviour or condition of a particular sub-group without the necessity for studying them again after one or more successive time periods. It is now the most

common methodology used to study changes in behaviour or attitudes, biological and cognitive effects of human ageing and social, political and cultural change. Cohort analyses in developmental and ageing studies are distinguished from longitudinal analyses, in which the same birth cohorts are re-examined at intervals over a period of time. A compromise is cross-sequential analyses, in which different age groups and birth cohorts are repeatedly re-examined and compared with themselves and with others at different measurement points.

Unfortunately, although cohort analyses always allow us to identify and study each of these effects, they do not provide any direct way of examining all of them, independently, in a single study. One reason for this is sampling variability. That is, any cohort samples we can obtain and compare are unlikely to be precisely comparable. A related difficulty, which usually guarantees that samples in gerontological comparisons will not be comparable, is sample attrition. As cohorts age, they lose members and so alter in terms of their credibility as representative samples of the populations from which they were initially selected. The most important limitation on cohort analyses is that there is no way to avoid confounding at least two of the three variables in which we are usually interested: age, cohort and period. This imposes very inconvenient restrictions on statistical analysis; for example, in a multivariate regression analysis, all three variables cannot simultaneously be entered as variables in a regression equation. Although many attempts have been made, and much has been written on the subject, there are still no statistical methods that can clearly separate the effects of birth cohort and period from the effects of ageing. Perhaps the most lucid and helpful discussions have been by Costa and McCrae¹ and Palmore². The current consensus is that decisions must rest on scientific judgement as to which two of the three possible effects are likely to be important, so that the third can be omitted. The

obligation to carry out exploratory data analyses to ensure that the effect that is to be neglected is in fact, statistically unimportant, should not need emphasis.

The inevitable confound between age, cohort and period effects is awkward because it means that we cannot use this most economical of all research strategies to solve all of the many problems connected with ageing. However, bearing their overriding limitation in mind, it must be stressed that cohort analyses are not merely an economical, but methodologically flawed, substitute for laborious longitudinal studies. They rather provide a means by which we can answer classes of questions that cannot be approached in any other way. The fact that the cohorts need not contain the same individuals is a statistical advantage as well as logistical convenience. For example, it can be very useful to sample quite different populations of the same birth cohort at different times in their lives so as to make an intracohort trend study. There are also unique advantages in comparing n quite different age samples of a population recruited at successive periods of time. This allows us to draw up what has become known as a "standard cohort table", in which the different age cohorts are ordered in rows, and the successive years in which these different age groups were sampled are ordered in columns. This allows us, at least, a rapid and convenient way to carry out exploratory analyses of our data. We can pick out intracohort trends, which become apparent by scanning down the diagonals; that is, we can see how members of a particular age group (whether represented by the same individuals followed over time,

or by quite different samples) are affected by both the passage of time (and so, among other factors, by their own biological ageing) and also by changes in the social, economic or epidemiological contexts in which they were studied. Within each of these sampling periods we can compare age cohort differences by scanning down the columns. Finally, by scanning along the rows, we can examine period effects which occur as one age cohort replaces another.

No single technique of comparison, whether cohort analysis, longitudinal analysis or even cross-sequential analysis, can provide a universal methodological panacea. Rather, each can solve problems that the others cannot approach, and none save us the effort of carefully thinking through the questions that we wish to ask of our data, and intelligently considering the implications of the comparisons we must make to answer them. To study the effect of ageing we must simultaneously acquire both cross-sectional and multiple longitudinal data and interpret them as sensibly as we can.

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Chronological and Functional Ageing

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Chronological age is the major descriptor by which aging is defined. In an effort to characterize the dynamics and variability of the aging process, recent approaches have used measures of functional aging to reflect the observation that individuals may function at a level above or below that expected for their chronological age. The relationship between chronological and functional age has implications for major questions of aging research and public policy. Do humans have a fixed lifespan? How can we maximize functioning for any given age? How is functional aging associated with vulnerability to disease? Should retirement be mandatory at a certain chronological age or should functional age play a role in this decision?

CHRONOLOGICAL VS. FUNCTIONAL AGING

Chronological age is defined as time since birth; its effect is analyzed in virtually all health studies and indeed in nearly all areas of human research. Although chronological aging is strongly associated with mortality and nearly all diseases, it should not be viewed as an etiologic factor. It is rather a proxy for numerous factors that change or accumulate over time, such as cumulative exposure to toxins and trauma, changes in hormone levels, immunological defenses and genetic repair mechanisms. A great deal of work has gone into identifying factors that explain the relationship between chronological age and susceptibility to disease and dysfunction, but much of this relationship remains unexplained. Identifying chronological age as a risk factor may be of little value, not because age is not strongly related to disease incidence but because an intervention to change one's age is not available. For developing prevention strategies, however, understanding an age association may be important, such as the finding that bone loss accelerates for 10–15 years after menopause, slows for a number of years, and again accelerates after age 75.

A number of different concepts, including functional age, biological age and biomarkers of aging, have been developed based on the observation that physiological measures and functional performance show a range of values in a population of a given chronological age. Functional age is a concept that rests on the premise that a measure other than chronological age could better reflect one's position in the aging process. Although biological and functional age have been defined differently, they are frequently used interchangeably. In contrast to chronological aging, which occurs at a universal fixed rate, functional aging has been termed "non-chronological" because its rate may accelerate or decelerate and, in fact, functional age may be greater or less than one's chronological age.

The concept of biomarkers of aging is particularly appealing because it implies that there are biological measures that reflect the rate of aging and that successful interventions on the aging process would have an effect on these markers. It is unclear at this time whether there is an underlying biological state of aging that can be summed up as a single number that indicates how far along in the aging process the individual has progressed. It has been argued that aging is a complex, uncoordinated phenomenon that can not be summed up in such a way¹. Others believe that, although a well-validated set of biomarkers does not currently exist, there may be techniques developed that can work well as general indicators of biological age². The requirements of a biomarker are not that it simply be different in persons or animals of different ages but that, in a group of subjects of the same age, it has a distribution of values that relates to other age-sensitive traits, such as longevity².

Much of the work done on functional aging has focused on physiological changes that are part of normal human aging. Another aspect of functional aging, which may be termed "functional health status", assesses functioning at the level of the whole older person, describing how that person functions in daily life. Functional health status has been found to be related to chronological age, disease and a variety of other modifying factors. Measures of functional health status have also proved valuable for clinical and health services research³. The USA's national goal to increase the span of disability-free life exemplifies the high level of interest in the measurement of functional health in recent years⁴.

FUNCTION VS. DISEASE IN CHARACTERIZING OLDER PERSONS AND OLDER POPULATIONS

Understanding the functional aspects of aging has been an important part of geriatric medicine for several decades. In the medical model of disease, the clinician gathers symptoms and signs, makes a diagnosis, and bases the therapeutic approach on this diagnosis. Complementing this disease-orientated approach, functional assessment provides an understanding of the impact and consequences of the older person's disease or diseases, giving information on level of independence and prognosis, as well as health care, rehabilitation and social needs. As aging research has increased in recent years, the functional approach has played an important role in its agenda.

Although normal physiological changes with aging may have an impact on the older person, the far greater functional impact comes from the effects of disease. A framework that represents the

relationship between disease and disability is therefore valuable in developing the concept of functional aging. The World Health Organization has proposed a theoretical pathway progressing from disease to impairment to disability to handicap⁵. An alternative pathway, proposed by Nagi⁶ and utilized by the US Institute of Medicine⁷, progresses from diseases and conditions to impairment to functional limitation to disability. An effort to operationalize this latter pathway defines “impairment” as dysfunction and structural abnormalities in specific body systems, “functional limitation” as restriction in basic physical and mental actions such as ambulating, grasping and stepping up, and “disability” as difficulty in doing activities in daily life such as personal care, household management, job and hobbies⁸. Although these pathways have remained essentially theoretical, increasing efforts are now under way to use empirical data to document important steps along the pathway⁹.

Disease severity and the co-occurrence of multiple diseases (comorbidity) play an important role in the process of disablement. In a study using data representative of the older US population, it was found that the prevalence of disability increased with increasing number of chronic conditions, after adjusting for age and sex¹⁰. The synergistic effect of specific pairs of diseases on disability has also been demonstrated. It is also clear that intervening behavioral, environmental and social factors play an important part in modifying the pathway along its entire course and need to be understood more fully^{11,12}.

DOMAINS OF FUNCTIONING AND MEASURES OF POPULATION DISABILITY

Functional aging is a multidimensional concept for which several domains must be considered to adequately characterize the total older person. Domains central to aging include physical, cognitive, psychological, sensory and social functioning. In the older population it is of value to separate the cognitive and psychological domains, although at times it may be difficult to ascertain whether cognitive impairment is related to a dementing disease or depression. The importance of sensory impairments in limiting overall functioning is receiving increasing attention in ongoing gerontological research. Social functioning, critically important in the lives of older persons, reflects the impact of the other domains on interactions with family, friends and the community.

Physical functioning, to be the focus of the remainder of this chapter, has traditionally been assessed through self-report of the ability to perform specific tasks, including self-care activities such as bathing and dressing (activities of daily living) and activities necessary to maintain independence in the community, such as shopping and food preparation (instrumental activities of daily living)^{13–15}. In cases where individuals have severe physical or cognitive impairment, proxies have been successfully employed to assess functional status, although proxies who have limited contact with the subject tend to provide less valid information¹⁶. Recently, performance measures of functioning, in which the individual is asked to actually perform standardized tasks, have been employed¹⁷.

A number of national surveys have estimated the prevalence of functional disability in the US population¹⁸. The prevalence of disability in activities of daily living (ADLs) in the non-institutionalized population rises steeply with increasing age and is slightly higher for women compared with men at the older ages (Figure 13.1)¹⁹. These rates do not reflect that portion of the population residing in nursing homes, where it is estimated that over 90% of residents require help with ADLs²⁰. It is important to keep in mind that, although chronological age is strongly related to disability prevalence, many other factors, such as socio-economic status, have large impacts on disability that are independent of age¹².

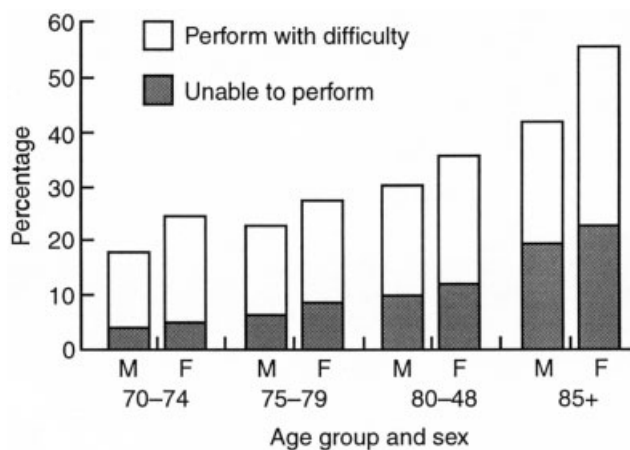


Figure 13.1. Percentage of persons who have difficulty or being unable to perform one or more activities of daily living by age and sex, USA, 1995. Activities of daily living include: bathing or showering, dressing, using the toilet, getting in and out of bed or chairs, and eating, walking and getting outside. Source of data: Second Supplement on Aging to the 1995 National Health Interview Survey

COMPRESSION OF MORBIDITY AND THE MEASUREMENT OF ACTIVE LIFE EXPECTANCY

An important issue related to functional aging is the relationship between length of life and the amount of time spent in the disabled state. Life expectancy has increased very substantially in this century. A consequence of this, which is just beginning to be appreciated, however, is that escaping death during the early years from infectious diseases and other causes may mean that many more people survive to ages where they suffer from chronic diseases, which can lead to long-term disability and loss of independence. A major goal of gerontology is to increase longevity without increasing the number of years spent in the disabled or dependent state. Although the recent increase in longevity is well documented, it is not now clear whether these added years of life have been accompanied by years of health and vigor or disease and disability. This question is of particular concern in the coming century, when it is projected that there will be continued increases in life expectancy and unprecedented numbers of old and very old persons. The theory of compression of morbidity predicts a future decrease in the number of years with severe disease and disability²¹.

An important tool for evaluating compression of morbidity is what has been termed “active life expectancy” or “disability-free life expectancy”²². Active life expectancy is defined as the average number of years an individual at a given age will survive and remain in the active, or non-disabled, state. Most analyses of active life expectancy have employed the ADLs to define disability, with active life expectancy calculated using life table techniques which consider transitions from the active, non-disabled state to both death and disability. The original analysis of active life expectancy considered the transitions to both death and disability as irreversible²².

However, recent longitudinal studies of aging populations have revealed that a substantial proportion of disabled persons make the transition back to the non-disabled state. Methods to calculate active life expectancy based on these kinds of changes, using multistate life tables, have been developed²³.

The relationship over time between life expectancy and active life expectancy can be used to assess the occurrence of a compression of morbidity. Three possible scenarios for population morbidity in women are illustrated schematically in Figure

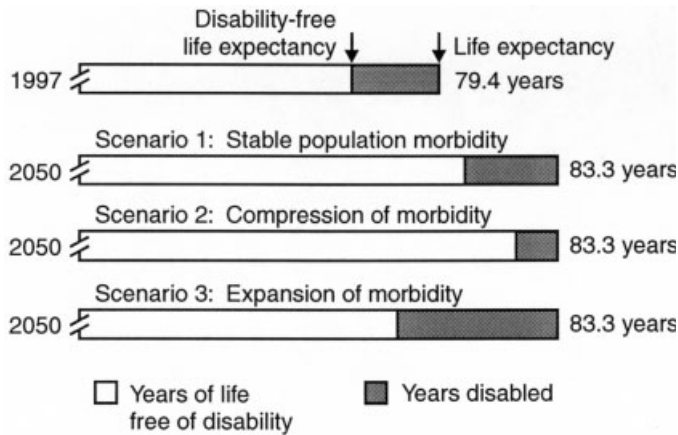


Figure 13.2. Scenarios for change in average burden of population disability level, 1997–2050. Compression of morbidity and alternatives

13.2. The total length of the bars in this figure represent life expectancy observed for 1997 and projected by the Census Bureau for 2050. The length of the unshaded segments of the bars represents active or disability-free life expectancy, and the shaded areas of the bars represent the average number of years in the disabled state. In scenario 1, the onset of disability has been postponed the same number of years as life expectancy has increased, and the number of years spent in the disabled state is unchanged from 1997. In scenario 2, there has been a compression of morbidity. Finally, in scenario 3, although disability-free life expectancy in 2050 has increased compared with 1997, it has not kept pace with increases in life expectancy and there is an expansion of population morbidity.

A vigorous debate over the prospects for a compression of morbidity began with a landmark paper by Fries, in which he made the claim that the compression of morbidity was inevitable in the coming years²¹. He argued that in all species the maximum lifespan is fixed, that human beings are quickly approaching this limit, and that with a stable life expectancy any postponement of disease and disability would result in a compression of morbidity. Although this logic is correct, others have pointed out that life expectancy is probably not going to reach its maximum level for at least the next half century and we must consider that any of the alternative scenarios depicted in Figure 13.2 are possible in the face of increasing life expectancy.

Repeat estimation of active and disabled life expectancy over time using identical techniques in the same target population would allow for a direct assessment of compression of morbidity, but these data are not available. However, disability prevalence, which is not equivalent to disabled life expectancy but reflects a cross-sectional picture of the proportion of the population that is disabled, can be estimated, and a great deal of attention has been focused on longitudinal trends in disability. In the 1970s there was some evidence of a rising prevalence of disability, but in the 1980s rates appear to have declined^{24–26}.

For example, Manton *et al.*^{24,27} have reported an average annual adjusted decline of 1.1% in the prevalence of a composite measure of severe disability in the population aged 65 and over, from the US National Long Term Care Survey for the period 1982–1994. Freedman and Martin²⁶ reported similar declines in a more specific measure of difficulty in walking or use of mobility aids from the Survey of Income and Program Participation: for example, the prevalence of disability defined by this criterion declined during the 10 year study period from

30.6% to 27.2% at age 65–79 and at age 80 and over from 44.0% to 40.9%. Although there is a scarcity of longitudinal data from other countries, similar findings have been reported²⁸ with only a few exceptions where rates have probably been stable.

Possible causes of these declines in disability prevalence include environmental changes making daily tasks easier⁸, more intensive use of assistive devices²⁹ and other social changes, including attitudes to being active in old age. In addition, the proportion of older people who have little education (a potent risk factor for disability) has declined^{26,30}. There is also evidence of declining prevalence of some medical conditions in old age, especially cardiovascular disease. Health risk avoidance and improved diagnostic and therapeutic techniques will also have contributed.

The future burden of morbidity and disability in the older population is of great concern to those involved in planning, financing and delivering health care and social services. If current rates of disabling diseases such as Alzheimer’s disease and hip fracture remain unchanged, the numbers of older people with these diseases will increase substantially in the next century³¹. Gaining an understanding of factors that have an impact on functional aging is critical if we are to reduce the burden of disability and achieve a compression of morbidity. Ultimately, effective interventions must be developed to prevent the onset and mitigate the consequences of diseases that lead to much of the disability in late life.

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Health Expectancy: Monitoring Changes in Population Health

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The increasing life expectancies experienced by many countries over the last decade have made the debate about the quality and quantity of years lived at older ages particularly relevant for conditions such as dementia, where the prevalence rises steeply with age. Health expectancies were developed to address the question of whether or not longer life is accompanied with a decrease in the quality of life, and they extend the concept of life expectancy to morbidity and disability by providing a means of dividing life expectancy into life spent in various states of good and bad health. Being independent of the size of populations and of their age structure, health expectancies thus allow direct comparison of the different groups that constitute populations: sexes, socioprofessional categories, regions.

As health expectancy combines a life expectancy with a health measure, there are as many possible health expectancies as health measures; for example, disability-free life expectancy, active life expectancy (based on independence in Activities of Daily Living), healthy life expectancy (based on good perceived health) or dementia-free life expectancy. Bone *et al.*¹ reports values for these and other health expectancies for the UK from two longitudinal studies of older people (the Melton Mowbray Ageing Project and the Nottingham Longitudinal Study of Activity and Aging) and from national cross-sectional studies. Dementia-free life expectancy at age 65 years has now been calculated for five countries: France, UK, Belgium, Eire and The Netherlands². Despite difference in life expectancies between countries (ranging for men from 13.5 years in Eire to 15.4 years in France and for

women from 16.9 years in Eire to 19.7 years in France), women can expect to live between one and two years and men between 0.5 and 0.7 years of their remaining life with dementia.

Today, estimates of health expectancy (generally disability-free life expectancy) are available for 49 countries³, although comparisons across time and between countries are still problematic due to the lack of harmonisation of measures and study designs. REVES (Réseau Espérance de Vie en Santé: the International Research Network on Health Expectancy) is an international organization of researchers, clinicians and health planners addressing these issues as well as developing and recommending methods of calculation and furthering the use of health expectancy as a tool for health planning.

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Health Expectancy: Monitoring Changes in Population Health

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The increasing life expectancies experienced by many countries over the last decade have made the debate about the quality and quantity of years lived at older ages particularly relevant for conditions such as dementia, where the prevalence rises steeply with age. Health expectancies were developed to address the question of whether or not longer life is accompanied with a decrease in the quality of life, and they extend the concept of life expectancy to morbidity and disability by providing a means of dividing life expectancy into life spent in various states of good and bad health. Being independent of the size of populations and of their age structure, health expectancies thus allow direct comparison of the different groups that constitute populations: sexes, socioprofessional categories, regions.

As health expectancy combines a life expectancy with a health measure, there are as many possible health expectancies as health measures; for example, disability-free life expectancy, active life expectancy (based on independence in Activities of Daily Living), healthy life expectancy (based on good perceived health) or dementia-free life expectancy. Bone *et al.*¹ reports values for these and other health expectancies for the UK from two longitudinal studies of older people (the Melton Mowbray Ageing Project and the Nottingham Longitudinal Study of Activity and Aging) and from national cross-sectional studies. Dementia-free life expectancy at age 65 years has now been calculated for five countries: France, UK, Belgium, Eire and The Netherlands². Despite difference in life expectancies between countries (ranging for men from 13.5 years in Eire to 15.4 years in France and for

women from 16.9 years in Eire to 19.7 years in France), women can expect to live between one and two years and men between 0.5 and 0.7 years of their remaining life with dementia.

Today, estimates of health expectancy (generally disability-free life expectancy) are available for 49 countries³, although comparisons across time and between countries are still problematic due to the lack of harmonisation of measures and study designs. REVES (Réseau Espérance de Vie en Santé: the International Research Network on Health Expectancy) is an international organization of researchers, clinicians and health planners addressing these issues as well as developing and recommending methods of calculation and furthering the use of health expectancy as a tool for health planning.

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Life Satisfaction

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Maddox and Wiley¹ suggest that the “relationship between aging and successful adaptation (variously ‘morale’ or ‘life satisfaction’ or ‘well-being’) is perhaps the most persistently investigated issue in the social scientific study of aging”. This statement, made a quarter century ago, remains true today. Life satisfaction is variously viewed as a prime indicator of overall life quality, a testimony to the importance of social structure and location for person well-being and one component of “positive mental health”². Life satisfaction research also has helped to balance the vastly overstated crisis orientation that characterized aging research in the past³. This chapter examines several facets of life satisfaction in later life: definitional issues, the epidemiology of life satisfaction in later life, major determinants of life satisfaction, mechanisms that may account for age differences in life satisfaction, and some final comments about future directions for life satisfaction research.

WHAT IS LIFE SATISFACTION?

Life satisfaction is most frequently defined as a global assessment of life quality, derived from comparison of one’s aspirations to the actual conditions of life^{4,5}. Because “life as a whole” or “life in general” is assessed, a long-range time perspective and non-specific life conditions are implied. Life satisfaction is one of a number of concepts that can be subsumed under the more general rubric of subjective well-being. Other indicators of subjective well-being are happiness and mood. Two primary factors underlie the distinctions among these concepts: the ratio of cognitive to affective judgment involved and time frame. Although all subjective well-being measures involve evaluation along a good–bad continuum, life satisfaction is primarily a cognitive assessment of the discrepancy between aspirations and achievements. In contrast, happiness and mood are primarily emotional judgments. In addition, happiness and mood are quite transitory, whereas life satisfaction tends to be stable (although it is sensitive to major events and changes in life conditions). Because it is a more cognitive, stable phenomenon, life satisfaction has been a more attractive candidate for the study of quality of life during old age. Researchers have understandably been more interested in relatively stable judgments of life quality than in indicators that fluctuate widely over brief periods of time.

How life satisfaction is defined has direct implications for methods that can be used to alleviate dissatisfaction. Dissatisfaction represents a significant discrepancy between one’s desired life conditions and one’s actual life conditions. Consequently, the two major ways to reduce dissatisfaction are either to engage in goal-directed behavior that brings achievements closer to aspirations or

to reduce aspirations so that they more closely match the actual conditions of life. As we will see, older adults are especially adept at using the latter strategy to sustain high levels of life satisfaction.

THE EPIDEMIOLOGY OF LIFE SATISFACTION

How satisfied are older adults with their lives? This question has been investigated for 40 years, with consistent results. Studies of samples of both older adults and of adults of all ages consistently document that the vast majority (i.e. typically 85%) are satisfied with their lives⁴⁻⁷. Between the 1960s and the 1980s, the life satisfaction of older Americans increased slightly relative to that of the non-elderly⁸. In studies conducted during the 1960s and earlier, older adults were somewhat less satisfied with their lives, on average, than their younger peers. By the mid-1970s, however, this pattern had reversed and has remained stable for more than two decades.

DETERMINANTS OF LIFE SATISFACTION

What are the major determinants of life satisfaction during old age? Research suggests that they fall into two primary categories: (a) objective life circumstances and (b) personality traits and other psychological characteristics.

Objective Life Circumstances

For all adults, regardless of age, the major determinants of life satisfaction include attachments to social structure (especially education, occupation and marital status), personal resources (including health and income), involvement in and support from primary groups (family and friends) and participation in meaningful social and leisure activities^{4,5,9-11}. Recent evidence indicates that religious participation also is a robust predictor of life satisfaction^{12,13}.

Although the same basic set of objective life circumstances underlie life satisfaction for all adults, the meaning and salience of those determinants may vary across ages or stages of the life course. What is highly important for a sense of well-being at one life stage may be less relevant at another. For example, George, Okun, and Landerman⁹ found that: (a) marital status is less important for a sense of well-being among young adults than among middle-aged and older adults; (b) income is most important for middle-aged adults and least important for young adults, and intermediate for older adults; (c) health is much more

important for older adults than for middle-aged and younger adults; and (d) social relationships are more important for younger and older adults than for the middle-age⁹. The degree to which the determinants of life satisfaction differ across age groups merits further investigation.

Despite evidence that life satisfaction is strongly related to objective life conditions, levels of life satisfaction typically remain stable, even in the face of substantial change in its documented predictors. It is true that significant social losses, especially widowhood and the onset of disability, are associated with decreases in life satisfaction. More surprising, however, is the fact that levels of life satisfaction remain stable in the face of other major changes in life circumstances. For example, although retirement typically results in a 50% reduction in income, there is no evidence that retirement and its accompanying reduced income lessens life satisfaction¹⁴. Indeed, life satisfaction can be sustained in the midst of potentially devastating chronic stress, as documented in a study of more than 500 family caregivers of demented older adults. Among this sample of caregivers, the average level of psychiatric symptoms was eight times greater than that for random community samples, more than one-third of the caregivers reported using psychotropic drugs, and nearly one-quarter used alcohol on a regular basis to cope. Nonetheless, fully 80% of these caregivers reported that they were satisfied or very satisfied with their lives—a figure only 5% lower than that typically reported in random samples of American adults¹⁵.

These findings testify to the distinctive character of life satisfaction—by definition, life satisfaction refers to subjective perceptions that go beyond the effects of objective life conditions. We need to know more about the cognitive and emotional processes that lead some older people who appear well off in terms of their objective life circumstances to nonetheless find life dissatisfying and burdensome. Similarly, we need to know why it is that many older adults who are less well off than their younger peers—and, indeed, than they were at younger ages—nonetheless find life meaningful and satisfying¹⁶.

Psychological Characteristics

One reason that life satisfaction may be overwhelmingly stable and only partially reflect objective life conditions may be that it reflects, in part, psychological characteristics. Several relatively stable personality traits are significant predictors of life satisfaction: neuroticism, extroversion vs. introversion, and openness to new experience or cognitive flexibility^{17–19}. Other social psychological characteristics are related to subjective well-being as well. For example, interpersonal trust is associated with higher life satisfaction²⁰. Personal coping strategies also have implications for life satisfaction. Selective optimization with compensation is a coping strategy in which older adults who face limitations due to health problems or diminishing social networks choose to increase their commitments to specific life domains in order to compensate for the inability to invest in other life domains. This strategy for coping with age-related decrements is associated with higher levels of life satisfaction than other coping strategies during late life²¹.

EXPLANATORY MECHANISMS

Before examining potential explanatory mechanisms that underlie subjective perceptions of life quality, another age difference in reports of life satisfaction must be noted. There is considerable evidence that older adults are satisfied with life conditions that are objectively worse than those needed to produce perceptions of equivalent life quality among middle-aged and younger adults^{8,9}. For example, some economists have observed that older persons

often express satisfaction with levels of income that are substantially below those required for minimally adequate food, shelter and medical care^{22,23}. Three types of explanatory mechanisms have been suggested as possible explanations for this pattern.

One possible explanation is aspiration theory which, as previously described, has been the basis of most conceptualizations of life satisfaction. From this perspective, some scholars argue that older adults are masters of the art of lowering aspirations to meet realities. A related hypothesis suggests that, because measures of life satisfaction focus on the long term, older persons often view current deprivations as piling in comparison to the dominant patterns of their lives^{24,25}. Although aspiration theory can describe plausible ways in which older adults can make peace with less than optimal life conditions, it cannot explain why older adults are willing to lower their aspirations substantially more than younger and middle-aged adults.

Second, other authors have suggested that younger adults use aspiration–achievement comparisons in making judgments about life satisfaction, whereas older persons assess the quality of their lives in terms of perceived equity. Research by Carp and Carp²⁶ suggests that young and middle-aged adults are satisfied when their achievements closely match their aspirations. In contrast, older adults are satisfied with life when they perceive their life conditions as fair and just (i.e. as matching what they “deserve”).

Third, and similar to equity theory, is the theory of relative deprivation, which posits that people will feel deprived only when they see themselves as worse off than others to whom they can appropriately compare themselves²⁷. There is strong evidence that most older people see themselves as better off than the “average older person”. According to relative deprivation theory, this kind of social comparison will result in high satisfaction. Equity theory and relative deprivation theory share the same problem noted above for aspiration theory. Although each can provide a plausible explanation for why older adults are often satisfied with less than their younger peers, neither explains why many older adults use these strategies to sustain high life satisfaction in the absence of optimal life conditions and most young and middle-aged adults do not.

FINAL THOUGHTS

Much has been learned from a half century of research on life satisfaction. We know and can rejoice in the fact that most older adults find life to be highly satisfying—as do most middle-aged and younger adults. We know that there are robust relationships between objective life conditions and perceptions of life quality. To the extent that this is true, high levels of life satisfaction among older adults confirm that older persons in modern societies share with their younger peers the material and social resources that make life enjoyable and satisfying. We also know, however, that older persons often express satisfaction with levels of resources that are below those necessary for quality of life. In this sense, we cannot deny the possibility that older people are not as well off in important ways as their subjective perceptions would suggest. This possibility—this nagging doubt as to the meaning of older persons’ reports of life satisfaction—raises two additional questions that merit increased scrutiny.

First, from a policy perspective, it is prudent to examine the objective life conditions of older adults’ lives as well as their ratings of life quality. We do not want, for example, to set standards for the economic resources needed to sustain quality of life on the basis of the thresholds that older people report if they, in fact, report levels that are below those needed to sustain adequate food, shelter and medical care. Similarly, we do not want to ignore the legitimate health and mental health needs of family caregivers of impaired older adults based only on the fact

that most caregivers report that their lives are satisfying. A related implication for policy analysis is the utility of life satisfaction measures for assessing the effects of intervention. Given the stability of life satisfaction measures in the face of all but the most major changes in life circumstances, it is probably unwise to evaluate the utility of an intervention on the basis of changes in life satisfaction. Such measures may be so insensitive that meaningful changes in life circumstances would be missed if life satisfaction were the only outcome examined. Moreover, most interventions need not improve life satisfaction to be demonstrably beneficial. For example, interventions that prevent illness or help to alleviate the functional consequences of illness need not improve life satisfaction to be beneficial. It is enough that they make a demonstrable difference in the incidence or functional consequences of disease.

Second, although it may be problematic for policy purposes that perceptions of life quality often stray rather far from objective life conditions, there is something admirable in the fact that most older people find life satisfying despite deprivations in the material and social resources so highly valued in modern societies. We need to investigate the foundations of this phenomenon. Increased understanding of this perplexing facet of life satisfaction is most likely to occur if subjective experience more broadly becomes the focus of investigation. In particular, increased attention should be devoted to studying the life of the self in old age and the sources of meaning that older adults attach to various life domains.

The 15% of the older population who are not able to sustain a sense of well-being also should not be ignored. Although we know that these older adults fare less well than their age peers on measures of income, health, social bonds and leisure participation, we know little else about them. We do not know, for example, whether their dissatisfaction in old age is a life-long pattern or is a response to age-related losses. Given the stability of life satisfaction measures, we also know little about how to facilitate higher levels of satisfaction among these individuals. Modern societies do quite well at providing the social and economic conditions that allow most older adults to find life satisfying. But we still do not know much about how to change life satisfaction. Such knowledge is of high priority, both for the practical purposes of promoting life quality and for gaining a better understanding of its dynamics.

Finally, we must remember that there is important information yet to be learned about life satisfaction. Life satisfaction is not the key barometer of successful aging (as, unfortunately, some of its advocates have implied). But it is important. It is important because it reminds the scientific community that we should be interested in enhancement as well as rehabilitation, in promoting the good life as well as intervening to alleviate bad times. Life satisfaction matters because it demonstrates that social, economic and psychological factors are as important in understanding life's triumphs as its tragedies. Life satisfaction is important because it reminds us to strive for health as well as the absence of disease and to invest in mental health as well as the amelioration of mental illness.

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The Normal Aged among Community-dwelling Elders in the UK

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INTRODUCTION

In 1851 there were approximately 1 million people in Britain aged 65 and over, representing 5% of the total population, compared with 9 million or approximately 16% in 1991 and a projected 11 955 000 in 2021¹. The reasons for this demographic transformation, which is not unique to the UK, have been discussed elsewhere in this book. One very visible manifestation of this change in the structure of our population is the increase in the number of centenarians. At the turn of the century, there were only a handful of British centenarian subjects, 200 at most². As we enter the next millennium there are approximately 6000 Britons aged 100 or more². One feature of population ageing in the next millennium will be the “ageing” of the ethnic minority populations of Britain, which will pose a new set of challenges³. Instead of this demographic trend being welcomed as a triumph for public health, population ageing is portrayed as an impending social disaster. Increasingly over the course of this century, old age has been depicted as a “social problem”, especially with regard to the health and financial consequences of an ageing population^{4,5}. Consequently, research in social gerontology has been dominated by the policy-makers’ agenda, which perceived old age as a social problem. The bulk of research has, therefore, concentrated upon defining and enumerating the social problem and pathological aspects of ageing, rather than on enhancing our understanding of “normal” ageing⁶. This is particularly true in the area of health and medicine where, until recently, dementia was seen as a part of “normal” ageing and where there are major debates as to how we are to afford to pay for the health care of older people. It is only recently that researchers have extended the horizons of their research to include research into the non-pathological aspects of later life.

One reason why the prospect of an ageing population is perceived so negatively is because of the prevailing images of the characteristics of the old age and older people generally. These images, beliefs and stereotypes are then transferred to the whole population. Stereotypes about the nature and characteristics of old age and older people abound and reflect a societal ideology which is predominantly youth-centred. To paraphrase the most common beliefs about older age, older people are seen as socially isolated, abandoned by their families, ridden with ill-health and disease (indeed, old age is synonymous with ill-health and

disease), consuming excessive amounts of societies’ health and social care resources and not contributing in any way to society (i.e. being parasitic upon the young, vigorous working population). Older people are further viewed as being inflexible, lacking vigour, being incapable of learning and shunning innovation and new technology. By inference, an ageing population is seen as being stagnant, uncompetitive and unable to compete or innovate as well as “young” populations. In the rest of this chapter we consider how valid some of these stereotypical images are.

THE LIVING ARRANGEMENTS OF OLDER PEOPLE

One very prevalent image of later life is that of living in institutional care. It is popularly believed that most older people live “in care” (either in a residential or nursing home). Although only approximately 5% of those aged 65+ live in some form of institutional care, this percentage is widely perceived as being much greater⁷. Even for the “oldest old”, those aged 90 or more, approximately half are still living in their own homes in the community⁸. For those older people living in the community, another common image is the association of later life with residence in “special” types of housing, such as sheltered housing. Again this is a type of accommodation characteristic of only a minority of older people, with approximately 10% of those aged 75 and over living in such types of housing¹. Hence, we can conclude that the majority of older people, some 70–99% (depending upon age) live in the community and in “ordinary” housing in the same way as the rest of the population. Home ownership is the main form of tenure, with 68% of those aged 65 and over owning their own homes in 1996/97¹. A further 27% lived in accommodation rented from the public sector¹. However across the tenure groups, older people are more likely than other age groups to live in older housing, which typically is in poorer repair, lacking in standard amenities and adequate heating. This is especially true for the minority of older people, approximately 10%, who live in private rented accommodation.

Within Britain, the post-war period has witnessed profound and far-reaching changes in patterns of family formation and household structure. In Britain in 1999, approximately 38% of those aged 65+ lived alone compared with about 10% in 1945⁹. Over the same period, the percentage of older people living with their spouse only has increased to about 50% from 30%. There

are, of course, important variations between men and women in terms of living arrangements. At all ages, because of the differential age at marriage between men and women and higher male mortality rates, there are more women living alone than men. For example, 26% of men aged 85 and over live alone, compared with 49% of women of the same age¹. Simultaneously, the percentage of older people living in larger households of two (or more) generations has decreased to about 10% of those aged 65 and over. Superficially the increase in "solo living" and decrease in the multi-generational household is often interpreted as evidence of the "rejection" of older people by their children and other relatives. However, these changes in household composition are, in fact, the result of much wider social changes, including the trend for younger people to establish independent households on or before marriage, geographical mobility, smaller families and increased rates of marriage combined with the improved financial resources available to older people, which allow them to express a widespread desire for independent living in later life.

SOCIAL NETWORKS AND SOCIAL CONTACT

Older people are often portrayed as a socially isolated group who are neglected by both their families and the wider social world. A key factor in considering the social networks of older people is the availability of kin and especially the marital relationship. The cohort of older women who never married, the post-World War I "spinster generation", has almost died out. Indeed, the percentage of single (never married) women has decreased from 16% of those over 80 in 1972 to 10% in 1999¹. For the current population of older people, approximately 7% have never married¹. The percentage of older people who are married varies markedly between men and women. Up to the age of 85 and over, the majority of men are married compared with 16% of women. For women the majority of those aged over 70 are widowed. Hence, for women the loss of a spouse can vastly restrict their social networks. However, the vast majority (97%) report the existence of close relatives and the presence of a close confiding relationship^{10,11}. Generally the pattern of contact between older people and their relatives and friends contradicts the image of neglect, as at least 75% of older people report weekly contact with relatives/friends¹. Amongst older women especially, friendship levels are both extensive and important^{12,13}. Furthermore, research has consistently demonstrated that only a minority of older people report that they feel lonely, although this percentage does increase with age and disability¹⁴ but is not synonymous with living alone¹⁴. Older people are also engaged members of society, as measured by their participation in social, cultural and voluntary activities¹.

HEALTH STATUS

Perhaps the most potent stereotype of all concerning later life relates to the perceived universal prevalence of illness, disability and infirmity. It is widely assumed that chronic ill-health, incontinence and mental impairment is the norm amongst the older age groups. How accurate is this image?

There are a variety of different ways of describing the health status of any given population. One widely used index of health status is mortality data. In Britain, mortality rates show a J-shaped distribution, being high in the first year of life and then decreasing in childhood. Rates then increase with age, so that 75% of deaths in England and Wales of the approximately 500,000 deaths annually were accounted for by those aged 65+¹⁵. In both Britain and the USA there have been significant reductions in later life mortality rates over the past four decades, which have contributed to increased life expectancy in later life. In 1980, men and women

aged 85 could expect to live, on average, for another 4.3 and 5.8 years, respectively. By 1994 this had increased to 5.3 years and 6.8 years because of decreases in later life mortality rates¹.

Whilst mortality data are useful, they tell us nothing about the health status of those who have not died. However, morbidity data are difficult to collect on a routine basis. One important British source of such data is the General Household Survey, which provides data about the prevalence of acute and chronic health problems amongst the general population of older people resident in the community. For acute self-limiting conditions prevalence rates increase only slightly with age, from 10% for those aged 16-44 to 20% at age 85+¹⁵. Chronic health problems, as measured by long-standing limiting illness, show a much stronger age gradient, increasing from 12% of those aged 16-44 to 53% at age 80+-91¹⁵. Although there are variations in disability prevalence estimates between studies, it is clear that chronic health problems are not universal in later life. Again, the last decade has, in Britain, been characterized by an increase in the number of years an older person can expect to live "free from disability"¹. Hence, the increase in life expectancy has not been accompanied by an increase in the amount of disability experienced by the "average" older person.

Within the elderly population there are important differences in health status, which largely mirror those of the population of working age¹⁵. For chronic ill-health, women show higher prevalence rates than men of the same age. For example, at 80 and over approximately 48% of men and 55% of women report that they have a long-standing limiting illness¹. Furthermore, there are significant differences in health status between the social classes. At all ages older people from professional and managerial occupations show rates of chronic ill-health which are significantly lower than their counterparts from the manual occupation groups¹⁶. Expectation of life at age 65 for men also varies from 16.8 years for those from social class I (professional occupations) to 12.6 years for those from unskilled occupations¹. These differences represent the continuation into later life of the well-documented social class differences in health status observed amongst the pre-retirement age groups.

There are significant methodological problems involved in quantifying the prevalence of dementia and other mental health problems in community samples. The varying methodologies and samples used often account for the varying prevalence rates produced by different studies. For both depression and anxiety there appears to be little significant increase with age. Depressive illness of clinical significance is observed in about 10% of those aged 65+, whilst the prevalence of anxiety is about 11% in those aged 65+¹⁵. For both conditions prevalence rates are higher in women than men. Dementia prevalence and incidence increase exponentially with age, regardless of the instrument used or sample studied¹⁷. A meta-analysis of 27 studies suggested that dementia prevalence increased from 1.4% at age 65-69 to 38.6% for those aged 90+¹⁷. However, even amongst the oldest age groups dementia is not universal.

For both chronic physical and mental impairment there are few data about how prevalence rates may be changing over time. Hence, in making comments about the health status of future cohorts of older people, we are extrapolating current prevalence rates. This may (or may not) be justifiable, as the health status of future cohorts of older people may be improved as a result of better living standards and access to medical care. There are two major stances concerning future patterns of mortality. Fries¹⁹ argues that morbidity will be compressed into the last few years of life, whilst the competing argument suggests that future populations will show increased rates of disability as more and more "frail" people survive into advanced old age²⁰. Whilst we cannot be certain if prevalence rates for chronic physical and mental health problems will change, and in what direction, in the

future, it is important to remember the limitations inherent in applying current health prevalence rates to succeeding generations.

POVERTY AND SOCIAL INEQUALITY

In contrast to the traditional views of later life as a time of poverty and deprivation¹⁹, there has recently emerged the image of the "Well-Off Older Person (WOOPIE)". This image has portrayed older people as a newly affluent group who are benefiting at the expense of younger people. However, recent studies have illustrated that, for the majority of older people, poverty remains the norm, especially in advanced old age¹⁹. In Britain, one in three of those living in poverty are older people and rates of poverty increase with age, are higher amongst women than men and amongst those from the manual occupation groups¹⁹. The explanation of such high rates of poverty amongst older people is the low level of state pensions, the principal source of older people's income, and the lack of access by many older people to additional sources of income, such as occupational and private pensions²⁰⁻²².

PATTERNS OF FORMAL AND INFORMAL CARE

Older people are the main users of the formal health and social care services provided in Britain⁹. However, despite this the majority of older people do not regularly use formal care services. For example, 19% of those aged 65+ use hospital inpatient services annually, 6% are visited by a district nurse and 4% by a home help⁹. Even for the potentially most vulnerable (those aged 85+ living alone), the district nurse/health visitor is in contact with less than 15%⁹. For most daily tasks that older people (and indeed other age groups) need help with, the principal source of support comes from the informal sector, usually either a spouse or children²³. Hence the family is, as it always has been, the major provider of care to the frail older person²³. However, we still know little about the ties within families that promote the development of the caring relationship²⁴. There is comprehensive research evidence which clearly shows that older people are themselves significant providers of care to each other²³ and younger age groups²³.

CONCLUSION

The research evidence now available illustrates that the common stereotypes of later life are both inaccurate and out of date. The majority of older people live in the community, in their own homes, and are integrated within a network of family and social relationships. The health status of older people is considerably better than the popular stereotypes, with chronic physical and mental impairment far from universal, even in the very oldest age groups.

Furthermore, although older people do use the health and social care services, the family remains their principal source of care. Indeed, older people are major contributors to the provision of care.

Moreover, within this less pessimistic review of later life, it is important not to treat older people as a single homogeneous social group. The experience of later life is greatly influenced by important dimensions of social stratification, such as gender, class and, increasingly in the future, ethnicity. It seems likely that such socially differentiating dimensions, especially ethnicity, will become increasingly important in determining the experience of future cohorts of older people.

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Do Life Events Seem Less Stressful to the Old?

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INTRODUCTION

Problems in defining, measuring and interpreting the impact of a life event have been much discussed^{1,2}. Where events occurring to older people are concerned, problems are compounded in that typically, mental health researchers assessing event severity are younger than the people assessed. A "social clock" view of development, however, assumes that the impact of an event depends on whether it is seen as normative or non-normative *for the stage of life*. The paradox is that older people are generally not the group that make the judgements of the severity of events occurring to older adults. A study was therefore carried out to see whether an elderly panel of raters accorded in judgement with a younger panel of four professional raters trained in the Bedford College LEDS scale, and whose ratings had been shown to be consistent and reliable³.

METHODS

A group of 25 retired university and health service personnel was recruited to rate the severity of the events and difficulties that occurred to a sample of elderly rural people over a 12 month period. The present results refer to the ratings made by a randomly assigned subgroup of 10 (age range 65–82) who were thoroughly trained in the LEDS procedures. Their ratings were compared with those of a (non-elderly) psychologists' rating panel. After training, the panel met for six sessions over a 3 month period to rate 80 life events drawn from a pool of 289 events, which had occurred to an independent sample of 237 rural elderly people living in the community. The events were drawn in a counter-balanced way according to the consensus severity (threat) assigned to the professional panel (I = severely threatening, IV = little or no threat) within each of 20 blocks. Training for the older panel used definitions and guidelines from the Bedford College Manual, together with taped interviews, group discussion and rating practice with feedback about the professional panel's ratings. After all blocks had been rated, five blocks were re-rated to examine reliability (20 events). There was a minimum of 1 week and a maximum of 3 weeks between ratings. All but one participant showed significant agreement between the two ratings (as measured by Kendall's coefficient of concordance). The inconsistent participant's data were excluded from further analysis.

RESULTS

The distribution of severity ratings for the older trained panel over the four levels of event severity were as follows: I, 3; II, 27; III, 34; IV, 16. A one-way analysis of variance (ANOVA) comparing the mean ratings of the old trained panel with those of the professional panel showed that there were significant differences ($F=5.6$, $P=0.004$). *Post hoc* comparisons indicated that the old group rated events as less severe than the young group. Cross-tabulations showed that, although there was some agreement of the older and professional panels (weighted kappa=0.31), the agreement with the professional panel was mainly about which events posed little threat. There was poor agreement for the crucial events of type I (severe threat). Of the 20 "severe threat" events presented, the distribution of ratings for the older panel were: I, 3; II, 6; III, 10; IV, 1. Most of these events concerned deaths of family members which, in general, the older panel (even though they had been trained in the LEDS procedures) did not see as particularly threatening.

CONCLUSION

The results provide support for the view that events such as deaths and severe illness events may be judged less threatening to older people, perhaps because they are seen as "on-time"⁴. Not all death events were rated equally, however. "Death of a spouse" and especially "death of a child" were regarded as more threatening than the death of a sibling or friend. However, in general the older panel rated stressful life events more conservatively.

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Support Networks

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It is difficult to find any text in social gerontology which does not refer to social networks, care networks or support networks. In the 1980s, a shift in social gerontology research took place to focus on networks of support rather than whether or not elderly people have contact with each of a range of categories of relationship (spouse, daughter, brother, etc.)¹⁻⁶.

Support networks here are defined as those within the larger social network of the individual who regularly provide support in a range of contexts of day-to-day life, and include: members of the same household; relatives seen most frequently; confidant(s); and those people providing, or perceived by elderly respondents to be available to provide, emotional support, instrumental help, personal care or advice. Support networks may range in size from two to 22 but modally cluster around five to seven^{4,7,8}.

Women have been shown to have more expansive networks than men in a wide range of countries⁹. Research has shown that older people with strong social networks are happier and more likely to perceive themselves to be healthy¹⁰. Others have suggested that, for people with serious mental illness, social support promotes normality in life styles¹¹. A review of the findings on social networks of older people⁹ makes the following observations, which have significance for geriatric psychiatry. Dense networks (where a high proportion of members know one another) tend to provide better access to emotional support but may shield or prevent people from seeking professional advice. For example, Veiel¹² found that for some patients, psychological and emotional support in crises received from close relatives was associated with a subsequent increase in depressive symptoms. Loose-knit networks provide better access to resources, including professional care. Residential admissions are less common where there is a close supportive network but this can also be associated with the avoidance of needed professional interventions.

The radius of support networks tends to follow a bimodal distribution, with substantial numbers of older people having either all members within 5 miles or at least one member more than 25 miles away. Support networks typically have a core of family but also include friends, neighbours and home helps. Normative expectations of support for different relationships exist that are well defined but these are likely to be culturally specific. Such expectations are hierarchical. In the UK spouses top the hierarchy, followed by immediate family: daughters/sons and sisters/brothers. Following siblings come friends, and then neighbours. Extended family of grandchildren, nieces/nephews and cousins come below friends and neighbours^{13,14}. Expectations are affected by the intervening variables of distance, gender and health. Long-term residents in a community tend to rely on local kin, while retirement migrants are unlikely to have kin nearby and have more diffuse networks^{15,16}.

The review of the literature referred to above suggests that older people may use different network members in different ways in emergency and non-emergency situations. There is evidence to show that vulnerable or stigmatized groups may be disadvantaged in network terms. Mentally ill people have smaller networks than others and the same is true for dementia sufferers. The more intimate (e.g. bodily care), personal (e.g. washing clothes) or private (e.g. financial concerns) the need of the elderly person (as defined by the culture), the more likely it is that this will be met by

a family member. However, non-kin appear to be more important than family for morale, self-esteem and emotional support.^{17,18} A high proportion of members of the support networks of elderly people are themselves elderly and/or female. Those who are or have been married tend to concentrate most need for support on one member of their network (usually a spouse or adult child), while those who have not married tend to spread their needs throughout their networks, relying on a wide range of others for one or two types of support.

Based on longitudinal data from the UK⁵, it has been shown that, contrary to expectations for largely elderly networks, the size of networks typically remains stable over time, suggesting some form of homeostasis. One exception to this is the support networks of married men, which tend to shrink with widowhood. The average change is equivalent to $\pm 1-2$ over 4 years. However, with increasing physical or mental frailty, networks tend to shift to reflect more reliance on proximate kin, if available, or increasing use of formal services and/or growing social isolation¹⁹.

Loss from networks is predominantly due to death or disability and gains come from the pre-existing social network, with members moving into the supportive core. Tensions in networks may arise from mismatch of expectations between the elderly person and the network member, i.e. where demands exceed normal expectations of the relationship or where different actors define the relationship differently. Mental illness, particularly the resultant cognitive dissonance experienced by network members, can result in network contraction as non-kin withdraw and kin tend to insulate the sufferer from non-kin contacts¹⁹.

Several different typologies of the networks of older people have been developed in Australia⁸, the USA²⁰, the UK⁶ and Israel²¹. What is striking is the similarity between them. Each typology: (a) reflects a continuum from close-knit to loose-knit; (b) finds that density is related to the importance of kinship; (c) includes the identification of a household-focused adaptation, based on a privatized life style and a small network; (d) identifies an association with social class (middle-class networks are less dense); and (e) acknowledges the influence of neighbourhood type.

Support networks demonstrate a range of types, which vary in terms of: availability of local kin; levels of interaction with different categories of membership; and the degree of community involvement, as measured by voluntary association. This article focuses on the Wenger support network typology, developed in the UK and subsequently validated in other countries. The Wenger typology identifies five types of network⁶, which have distinct parallels with the other network typologies. The five types of networks identified are summarized below. The first three types are based on the presence of local kin; the other two types reflect the absence of local kin:

1. *The local family-dependent support network* has a primary focus on close local family ties, with few peripheral friends and neighbours; it is often based on a shared household with, or close to, an adult child, usually a daughter. Community involvement is generally low. These networks tend to be small and the elderly people are more likely to be widowed, older or in less than good health.

2. *The locally integrated support network* includes close relationships with local family, friends and neighbours. Many friends are also neighbours. It is usually based on long-term residence and active community involvement in church and voluntary organizations in the present or recent past. Networks tend to be larger on average than others.
3. *The local self-contained support network* typically has arm's-length relationships or infrequent contact with at least one relative living in the same or adjacent community, usually a sibling, niece or nephew. Childlessness is common. Reliance is focused on neighbours but respondents with this type of network adopt a household-focused life style and community involvement, if any, tends to be low key. Networks tend to be smaller than average.
4. *The wider community-focused support network* is typified by active relationships with distant relatives, usually children, and high salience of friends and neighbours. The distinction between friends and neighbours is maintained. Respondents with this type of network are generally involved in community or voluntary organizations. This type of network is frequently associated with retirement migration and is commonly a middle-class or skilled working-class adaptation. Networks are larger than average. Absence of local kin is typical.
5. *The private restricted support network* is typically associated with absence of local kin, other than in some cases a spouse, although a high proportion are married. Contact with neighbours is minimal; there are few nearby friends and a low level of community contacts or involvements. The type subsumes two sub-types: independent married couples and dependent elderly persons who have withdrawn or become isolated from local involvement. Networks are smaller than average.

As a result of different network configurations and differing normative expectations for members, some networks are more robust in terms of the provision of the informal support they can provide. Different types of networks react in different ways to the common problems of old age^{22,23}. Different network types have different strengths and weaknesses and nature of the risk is related to the type of network²⁴. Network types have also been demonstrated to relate to levels of use of domiciliary services and to types of presenting problems^{25,26}. Knowledge of network type can, therefore, be a useful tool in planning therapeutic interventions and a measurement instrument for use by practitioners has been developed^{22,24}. This makes it possible to identify support network type on the basis of eight questions.

Recent research has paid increasing attention to the significance of network type for professional practice. It has been suggested that to achieve "real improvements in the quality of life", the relationship environment of people with severe, long-term mental health problems needs to be explicitly addressed by practitioners²⁷. Other authors²⁸ make the same point in the context of bereavement. Network type has been shown to be strongly associated with depression but not anxiety²⁹.

High needs for intervention have been identified in the context of a firm diagnosis of dementia associated with networks that reflect low levels of social support³⁰. Admission to long-term care of older people with dementia occurs sooner in all network types other than the family-dependent network. In network types without local kin, the network tends to become private, restricted in the face of dementia³¹.

The distribution of network types has been shown to be related to neighbourhood or community. Typically, where population stability is high, those types of networks based on local kin and providing high levels of informal support are found. Where population movement is high, for example in areas where retirement migration is common, the proportion of more

vulnerable network types is greater^{13,15}. Because of the relationship with coping strategies, knowledge of the distribution of network types can be an important indicator for planning the mix of service provision, in the same way that the identification of the network of an individual can indicate the appropriate type of intervention and potential future outcomes.

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World Statistical Trends and Prospects

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The aging process relates to both individuals and the collective aggregates that they comprise, such as families and nations. The prolongation of human life has resulted in ever-increasing numbers of persons not only reaching the threshold ages defining old age, but living beyond. In most of the world's developed countries, at least four-fifths of the persons born today can expect to reach their 65th birthday and, of these survivors to old age, over 40% will reach 85. At the beginning of this century, only half as many (two-fifths) could expect to reach age 65, with only 14% of the survivors likely to live to age 85. The future will no doubt bring even further extension of life for greater numbers of individuals. A corollary of this development will be increases in the numbers of older persons in our populations, given the large size of future cohorts of persons reaching the older ages.

This remarkable achievement in extending life, largely a phenomenon of the twentieth century, has important consequences for the well-being of individuals and the societies in which they live. Most notably, we must be concerned with the implications of these trends on the life conditions of older persons and their impact on the social, economic, political and health institutions of societies. As many contributions to this volume emphasize, loss of intellectual functioning and the emergence of mental illness are not inevitable concomitants of aging. But the older population does have the highest frequency of cognitive impairments and disruptive behaviors of any age group and are least likely to be seen by mental health specialists¹. Moreover, these conditions also command high levels of care and support from family and community networks. An increased awareness of the demographic context, both present and future, is necessary for furthering our understanding and treatment of these phenomena.

In this chapter, we provide a statistical treatment of the diverse dimensions of this aging process, particularly population aging, and how it is likely to evolve in the years ahead. Although population aging is truly a world-wide experience, for this volume we focus on conditions within the so-called more developed region (MDR), which include North America, Japan, Europe, Australia and New Zealand. It is in this more developed region that population aging has progressed the most in terms of the proportions of older persons in the population. Moreover, the countries in this region also have experienced major changes in the composition of the aged population itself. The data reported in this chapter are derived from the most recent population projections (medium variant series) of the United Nations, as assessed in 1998².

GROWTH OF THE OLDER POPULATION

The older population—persons 65 years and over—is estimated at 171 million for the more developed region in 2000. They

represented 14.4% of the total population, about one in every seven persons. The number of older persons has increased by 45 million or 35% since 1980, compared with an increase of only 6% for the under-65 population. By the year 2050, the region's older population is projected to reach nearly 300 million, a numerical increase over the 50-year period of 128 million. As noted in Table 16.1 and Figure 16.1, the growth accelerates after the year 2010, as persons born in the post-World War II period reach the older ages, then subsides somewhat after 2040.

The growth rates for the older population over the total period greatly exceed those of the under-65 population of countries in the region. In fact, the younger population actually declines in number after the year 2010, as newly-aged cohorts advance into

Table 16.1 Population distribution by age group, more developed region, 1980–2050

	0–14	15–64	65–79	80+
1980	243	714	126	22
1990	236	768	144	31
2000	216	801	171	36
2010	196	820	192	50
2020	193	791	232	58
2030	187	749	274	72
2040	182	712	294	92
2050	177	679	299	102

Source: United Nations, *World Population Prospects: The 1998 Revision*, Vol II.

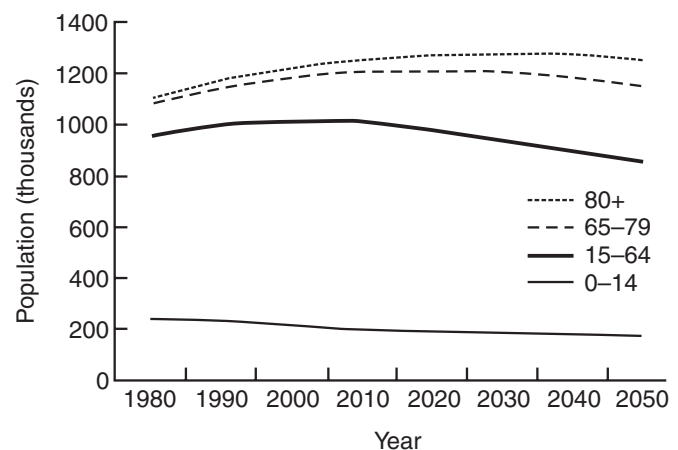


Figure 16.1 Population distribution by age group, more developed region 1980–2050. Source: United Nations, *World Population Prospects: The 1998 Revision*, Vol II

the older population and are not fully compensated for by youths. Thus, the proportion of older persons in the total population is expected to steadily increase, reaching nearly 23% in 2030. This aggregate figure masks considerable variation among countries, with Japan (27.3%), and Italy (29.1%) expected to reveal the highest levels and Ireland (17.9%) and New Zealand (18.8%) at the lower end. Nonetheless, the proportion of older persons for all countries in the region is historically unprecedented. An added dimension of these changes in the age composition of the populations is the expectation that by 2015 the proportion of older persons will exceed the proportion of young persons aged 0–14 years, given realization of the assumptions regarding fertility introduced into the projections.

AGE AND SEX COMPOSITIONS OF THE AGED POPULATION

Not only has the region experienced overall population aging, but the older population has itself aged. The growth of the oldest old (persons aged 80 and over) has important implications in terms of the health status of this group of people and their need for health care and social supports. If the assumption is correct that the prevalence of mental health problems increases with age, then these trends toward aging of the older population take on added importance. Moreover, the incidence of the most severe mental disorders (e.g. senile dementia of the Alzheimer's type) is greatly manifested at advanced ages³. The percentage of oldest old in the total older population showed a marked rise in the 1980s, when it increased from 17% to 22%. It will steadily rise until the year 2010 (26%) and then will decline slightly to 24% in 2025. The decline reflects the large number of persons who will reach the older ages between 2010 and 2030. Nonetheless, the overall numbers of oldest old will nearly double to 72 million in the region over the 2000 figure.

A noteworthy feature of the older population in the region is the preponderance of women over men. In 2000, there were 153 older women for every 100 men and, among the oldest old, 226 women for every 100 men. These levels for both groups are likely to be somewhat attenuated in the future, as the aged population gains large new cohorts, but the imbalance will still remain.

COMPONENTS OF POPULATION AGING

While the overall age and sex structure of a population is affected by fertility, mortality and migration, the main factors producing changes in the older population result from the relative size of cohorts reaching the older ages and the survival of these older persons to further advanced ages. Although the major source of future growth results from the large baby boom birth cohorts that will enter the older population, as was noted previously, the improved chances of persons reaching old age and surviving to advanced ages have also been a potent force in producing the present and future growth⁴.

The average life expectancy of a baby born between 1995 and 2000 and experiencing the age-specific mortality conditions of that period was 71.1 years for a boy and 78.7 for a girl⁵. To gain some perspective on these figures, the following levels were estimated for the period 1970–1975 and projected for future periods:

	Male	Female
1970–1975	67.6	74.7
2020–2025	75.5	81.8
2045–2050	78.2	84.2

There was an increase of 3.5 years for males and 4.0 for females in the 25-year period preceding 1995–2000 and gains of 4.4 and

3.1, respectively, projected for a similar time period to 2020–2025. The reversal in the projected survival by gender can be noted and this is carried over into the next period, when only minor life expectancy gains are forecasted. Aside from this questionable gender reversal, some also view the assumptions underlying the UN projections as rather conservative. In fact, a life expectancy at birth for females of over 84 years has already been attained in Japan and is likely to be exceeded in many other countries by the second decade of the twenty-first century.

The extension of life has meant that the remaining years of life after persons reach age 65 have increased markedly. For the USA, a surviving male could expect to live 15.7 years on average at age 65 in 1996 compared with 11.5 in 1900, a 36% increase⁶. The average female could expect to live an additional 19 years in 1996, a gain of 6.8 years or 55%. Moreover, 76% of men and 86% of women born in 1996 can expect to reach age 65, doubling the percentages in 1900. Thus, not only have more persons reached old age, but an increasing amount of one's life is being spent as an older person.

These improvements in survival have resulted from widespread reductions in communicable diseases and even declines in the lethal progressions of some of the predominant chronic, degenerative diseases. A noteworthy feature of overall mortality trends in the region has been the reductions in death rates at older ages. An emerging issue of considerable policy importance is the extent to which these improvements in longevity have been accompanied by increases in active life expectancy free of major chronic diseases, functional disabilities and institutionalization. While data are lacking to definitively establish past developments, there is evidence that active life expectancy has tended to increase, perhaps even keeping pace with improvements in overall life expectancy⁷. However, with increased population aging, the net result of these developments is the likelihood of increased numbers of persons at old ages with chronic co-morbidities and disabilities and, therefore, in need of long-term care and rehabilitative services.

SUPPORT STRUCTURES

The demographic parameters that relate most directly to these emerging issues concern the households and family conditions of older persons who are likely to need supportive assistance (care-receivers) and those who are potential care-providers. A number of complex demographic and social factors are involved in this matrix.

There has been a steady trend over time toward increased proportions of older persons being married and reduced proportions being single and widowed in the countries of this region⁸. Moreover, this development is true not only for the total aged population, but also for all age groups up to the most advanced. However, it should be noted that a much larger proportion of men than women are likely to be married, with the converse true of widowhood. These trends are a product of both the previous marital experience of persons in cohorts reaching the older ages and improvements in the joint survival of spouses. Nonetheless, there are strong signs that this pattern will change in the next century, as large cohorts of persons who are divorced or have never married enter into the older ages.

Another distinctive trend has been the increasing likelihood of older persons, both married couples and individuals, to live independently from other family members or non-relatives. This pattern of living arrangements reflects the smaller size of families resulting from prior declining fertility, as well as value preferences among older persons for independent living. However, these observations say little about the degree of contact among family

Table 16.2 Generational support ratios, more developed region, 1980–2050

	45–49/65–70	65–69/80+
1980	0.62	2
1990	0.58	1.56
2000	0.64	1.41
2010	0.64	1.1
2020	0.48	1.23
2030	0.41	1.08
2040	0.37	0.8
2050	0.35	0.71

Source: United Nations, *World Population Prospects: The 1998 Revision*, Vol II.

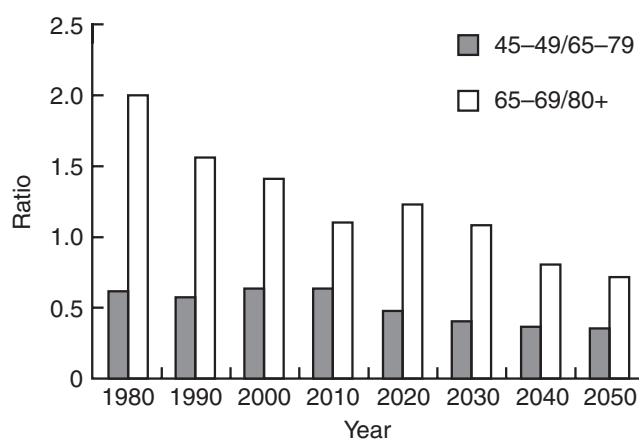


Figure 16.2 Generational support ratios, more developed region, 1980–2050. Source: United Nations, *World Population Prospects: The 1998 Revision*, Vol II

members or the strength of social networks, which appears to have remained strong.

Over the last few decades, increasing numbers of older persons have resided in institutions, although the proportions of such persons have not risen markedly in most developed countries⁹. While only 5–10% of older persons are in institutions at any given time, approximately 40% of persons will be institutionalized at some point in their lifetimes. Future prospects in light of the growing number of older persons living alone have given rise to considerable concern about the social, medical and economic issues of providing non-institutional vs. institutional care. Moreover, there is considerable policy debate over the optimal balance between family and state-provided care, whether in the home or in a more formal setting.

This brief review of trends suggests that there are major developments emerging from the growth of the older population and increased longevity that signal important changes in the potential care burden for vulnerable persons in the aged population. In turn, it raises questions about potential caregivers, most notably members of younger generations. A broad indication of the demographic indicators can be gained from examining ratios of the number of persons who are aged 45–49 to those aged 65–79. The former can be thought of as a group consisting of

children of older persons. A second ratio relates two generations of older persons—those aged 65–69 to those aged 80 and over.

Table 16.2 and figure 16.2 show these two sets of ratios over the period 1980–2050. The importance of these ratios is not in their absolute levels, but in the relative levels over time. The younger generational ratios reveal little meaningful change up to year 2010, but thereafter decline steadily as the baby boom generation enters the older ages. The older generation ratios show more fluctuation. They have declined sharply from 1980 to the present and will continue downward to 2010. They show a rise in 2020, but then slowly decline to below parity through the projection period. In general, the shorter-term trends are somewhat reassuring in terms of the availability of potential caregivers but the longer-term trends reveal the increasing pressures on support systems. It should be noted, however, that such ratios do not inform us about the capacity of younger generations to provide assistance, which can be affected by kin relationships and involvement in employment that precludes extensive assistance.

SUMMARY

Population aging is a pervasive and seemingly irreversible process for countries in the world's more developed region. Not only will older persons increase dramatically in number and as a percentage of total populations, but dynamic forces will modify the socio-demographic and health status composition of aged populations as well. Like the Struldbrugs or Immortals of Swift's *Gulliver's Travels* the advantages of life prolongation have complex consequences, not only for the individuals involved but also for the societies in which they are found.

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Demography of the Old: Implications of Recent Trends

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In most developed countries, at least 10% and in some cases more than 15% of the total population is aged 65 and over¹. Recent projections suggest that by 2025 the proportion aged 75 and over will lie in this range; by then people aged 65 and over will contribute a quarter of the total in a number of countries including Japan, Italy and Greece, and over one-fifth in most of the rest of Europe. The proportion of “oldest old”—those aged 85 years or more—is increasing particularly rapidly². As the risk of dementia is so strongly age-related, these changes will mean a large increase in the numbers affected unless strategies for prevention and postponement can be identified and implemented. However, there are also other implications of current demographic trends. Here I focus on changes in mortality and survivorship and also consider briefly changes in marriage.

The initial impetus for this ageing of the human world came from the transition to lower fertility that was set in motion in many Western populations in the last quarter of the nineteenth century and is now close to being a global phenomenon. More recently, the pace of population ageing, and the ageing of elderly populations, has been accelerated by marked declines in late age mortality; these are now the predominant motor driving the further ageing of populations with already old age structures³.

The extent of this change is illustrated for five developed countries in Table 1. In England and Wales, France and the USA,

male expectation of life at age 65 increased as much or more between 1970 and 1995 than in the preceding 70 years (or indeed, in the case of England and Wales, in the preceding century⁴). Among women too, the extent of improvement since 1970 is notable. Viewed from a cohort perspective, increases in longevity appear even more remarkable. Table 2 shows the proportion of each birth cohort surviving to the age of 80. Survival to later older age has increased dramatically reflecting not only large improvements in infant and child mortality (still very high in the nineteenth century) but also the more recent improvements in late adult death rates. Today, women of 80 represent not a select group of elite survivors, but *half* their original birth cohort. These improvements are not confined to the “young” elderly; death rates among the oldest old, including centenarians, have also dropped substantially^{5,6}.

These changes have sparked fierce debate about possible changes in the relationship between mortality and morbidity, the plasticity of the ageing process, possible limits to longevity and, of course, implications for health services. Less frequently considered are the implications for the family and support networks of older people and how these may interact with other changes in, for example, marriage and partnership.

The gender differences in mortality apparent in the tables underlie one of the most notable features of the older population, the preponderance of women. In the UK in 1998 the ratio of women to men at age 60–64 was close to parity (1.04) but rose to 1.85 at age 80–84 and 3.5 in the 90–94 year-old group. One consequence of this gender imbalance, compounded by the common pattern of women marrying men older than themselves, is that 65% of women aged 75 and over are widows, while 62% of men of this age are still married. If the narrowing of sex differentials in mortality continues, as is projected, the proportion of widows in this group will fall to 52% by 2021; however, by then 10% will be divorced (compared with 3% now) and a much larger proportion will have experienced divorce at some point in their lives (21% of those born in 1950 had already experienced divorce by the age of 40⁷). Those who have experienced marital disruption have poorer health in early old age⁸, although this may be because of factors associated with both risk of divorce and poor health.

Changes in the age structure of populations also mean change in the age structure of families. Lower mortality increases the availability and duration of “vertical” kin networks. A recent (1999) national survey showed that 74% of people aged 80 and over in Britain were members of three-generation families and over one-third had living children, grandchildren and great-grandchildren⁹. Contacts between older people and their children and grandchildren are frequent; however, there have been very large decreases in the extent of intergenerational co-residence. In 1991 21% of women aged 85 and over lived in two- or three-generational households, compared with 42% 20 years earlier¹⁰. During the same period (1971–1991) the proportion of older people moving to institutions increased, while the proportion moving to live with relatives fell, suggesting some substitution of the former for family care¹¹.

Demographic changes of the type and magnitude, which here I have had space to allude to only briefly, require some thinking about roles, relationships and activities throughout the life course.

Table 1 Further expectation of life at age 65, 1900–1995

	1900–1901	1950–1951	1970–1971	1995
Men				
England and Wales	10.1	10.8	11.9	14.8
France	10.0	12.2	13.0	16.1
Japan	10.1	10.9	12.5	16.5
Sweden	12.1	13.5	14.3	16.0
USA	11.4	12.8	13.1	15.3
Women				
England and Wales	11.1	13.4	15.8	18.4
France	10.9	14.6	16.8	20.6
Japan	11.4	13.0	15.4	20.9
Sweden	13.0	14.3	16.9	19.7
USA	12.0	15.1	17.1	19.2

Data from: Government Actuary's Department, Berkeley. Mortality Database (<http://demog.berkeley.edu/wilmoth/mortality>); and the Japanese Ministry of Health and Welfare.

Table 2 Survival to age 80 by birth cohort, England and Wales

Year of birth	Survival to age 80 (%)	
	Men	Women
1861	10	16
1881	14	25
1901	17	34
1921*	29	47

*Partly based on projections.

Source: Author's analysis of data from the Government Actuary's Department.

Most people now spend much longer as the parent of an adult child than they do raising minor children; increasing proportions will have step- and half-kin of various kinds and will live to know their great-grandchildren. However, expectations about these types of relationships may be confused. The increasing diversity of partnering and parenting patterns may also have implications both for mental health in later life and the support of those with mental health problems.

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Part C

Abnormal Ageing

The Influence of Social Factors on Mental Health

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Social factors have an enormous influence on the life course, affecting development and socialization, the relative influence of families and peer groups, opportunities for work, recreation and social participation, and patterns of social integration and independence as one reaches the later years. With the rapidity of social change, persons living a normal lifespan are required to modify their expectations and behavior on many occasions if they are to adapt successfully to shifting social conditions. Transformations in technology, sexuality, fertility, family and work life, household structure and many other facets of life, also make it inevitable that different age cohorts will have diverse life experiences.

Almost every aspect of mental health and well-being are influenced by social factors and social institutions¹. The effectiveness of social institutions, and the extent to which they build supportive relationships, coping capacities and personal commitment, contribute importantly to mental health outcomes. The dynamics of populations, and the distribution of persons at varying ages, also affect the productive capacities of nations, the prevalence of disability and dependence, and the capacity of a society to protect its citizens against the risks of disadvantage. Social attitudes and patterns of community organization can either encourage or inhibit full participation and meaningful roles for children, the elderly, the disabled or other groups, affecting functioning, quality of life and psychological well-being.

The discussion that follows focuses more on severe mental illness than on levels of psychological distress or emotional well-being. The distinction is not clear-cut because even depressive symptoms short of a clinically diagnosable disorder can have devastating effects on functioning and quality of life and are a major source of disability, surpassing in impact many serious chronic diseases².

Severe mental disorder, such as depression, bipolar conditions, and schizophrenia results from complex, but poorly understood, interactions between biological vulnerabilities and psychological and social influences. Adverse social and developmental factors may increase susceptibility to serious mental illness, or may contribute to triggering illness among vulnerable persons. Among persons with mental illness, social factors may significantly ameliorate symptoms, influence treatment patterns and enhance or impair quality of life. Most of the major social factors—age, gender, social class, race and ethnicity, familial arrangements and the like—are associated with mental illness, either by contributing to its onset or course or because of social selection factors. Moreover, there are complex interactions among social factors

such that, for example, the impact of socioeconomic status or gender on mental health outcomes may vary by age or by birth cohort.

The epidemiology of illnesses that first occur later in life may be quite different from chronic conditions that persist through much of the life course. Understanding the occurrence of mental illness later in life, however, is complicated by the relationships between physical illness, drug use (commonly prescription drugs) and the occurrence of symptoms consistent with mental illness. Prescription and non-prescription drug use is high in elderly populations because of the prevalence of illness. The inappropriate use of pharmaceuticals among elderly persons is also common and includes physicians over-prescribing to older patients, self-medication and drug interactions³. Moreover, drug sensitivity changes with age-related changes in individuals' capacity to absorb and metabolize drugs; therefore, dosages effective in younger patients may be ineffective or excessively high for older patients. There continues to be considerable concern that institutional settings over-use medication in order to sedate and control patients. Drug reactions, including confusion, hallucinations, paranoia and mania, are common and may be inappropriately diagnosed as mental illness.

Schizophrenia typically first occurs in late adolescence or in early adulthood and may have a complex and fluctuating course. Follow-up studies of early-onset schizophrenia indicate that, with aging, the positive symptoms of schizophrenia abate or remain in remission for longer periods of time, and that for a significant minority of persons complete remission is possible, although the processes that lead to such outcomes are not well understood⁴. Selective mortality or the natural course of the disorder may be responsible for the relatively positive outcomes observed among older persons with schizophrenia. It is also possible that lowered personal expectations, the development of coping strategies, improved adaptation and learning how to avoid upsetting stresses also contribute to positive outcomes for persons with schizophrenia. On the one hand, persons with chronic schizophrenia bring into their later years life histories that are likely to be characterized by significant periods of disorder and disruption. On the other hand, they bring an array of skills and coping strategies, developed through dealing with illness over many years, and these may mitigate the potentially negative impact of the illness during later life.

Late-onset schizophrenia is relatively rare and little is known about its course^{5,6}. The lack of evidence of late onset of schizophrenia and other psychoses among the elderly may in part result from diagnostic practices that give priority to co-occurring dementia and confusion that are highly prevalent in

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later years. It has only recently been recognized in the USA, and there have been very few community-based studies examining outcomes. There is some suggestion, however, that those persons with onset later in life are able to function better in the community because occupational and social roles have not been interrupted by the disorder earlier in life³.

There has been more research attention focused on depression during later life and although there have been widespread reports of a high prevalence among this population, epidemiological studies have not confirmed this impression⁷. But it is unclear whether the relatively low rates of depression at older ages are artifacts of epidemiological definitions and methods. Sub-threshold negative affect appears very high in this population, leading many internists and geriatricians who care for the elderly population to believe that epidemiological estimates are biased and do not reflect the real situation. Depression is clearly different in elderly persons than in early and middle adulthood because of complex physical co-morbidities, decrements in function and relationship losses that occur in later life. Older persons are also less likely to acknowledge psychological manifestations of depression, see mental health treatment as more stigmatizing and resist referral to mental health specialists. Assessing depression within the constellation of physical concomitants that might be attributed to a variety of other conditions poses diagnostic challenges for the general physician. There are presently a number of ongoing experimental efforts to improve the recognition and treatment of depression among the elderly in primary care⁸.

Most studies examining depression among elderly persons have not dated the onset of the disorder. There is some suggestion, however, that biological or genetic vulnerability may play a lesser role in the etiology of first onset of major depression or bipolar disorders late in life than earlier^{9,10}. Late-onset depression and bipolar disorder appear to be triggered more commonly by medical and neurological co-morbidities, while onset earlier in adulthood is more strongly correlated with the occurrence of environmental stressors¹⁰. Much remains to be learned about the etiology of late-onset disorders but the role of social factors will vary with age.

Recent research reports a gender difference in late-onset schizophrenia, with older women having greater risk⁵, although the reason for this difference remains unclear. There is also a gender difference in depression but, unlike schizophrenia, it develops in early adolescence and persists across the life course. The relationship is a complex one and is a matter of continuing controversy. Since abuse of alcohol and drugs and violence are more common among men, and depressed affect more common among women, some explain the two-fold prevalence difference in terms of varying gender-related reaction patterns¹¹. Others have argued that the higher rates of reported and diagnosed depressive illness in women result from a greater prevalence of more common and less serious symptoms of depression reported by women, such as sadness and crying, and their greater willingness to report affective symptoms¹². The issues remain unresolved, with different views depending in part on varying conceptions of the nature of depressive disorders.

Whatever the eventual resolution of these issues, the data do suggest that women in later life are at greater risk for symptoms of schizophrenia and depression than are their male counterparts. At the same time, older women are more likely than older men to be without a partner and they also experience greater economic disadvantage, both of which may exacerbate the negative consequences of mental disorder.

There is a long-established relationship between socioeconomic status and severe mental illness and decades of debate about the relative influence of causative vs. selection factors for explaining the higher prevalence of schizophrenia among those of lower socioeconomic status^{13,14}. The weight of the evidence supports the

view that the impairment associated with schizophrenia prevents upward mobility comparable to one's age cohort and loss of social position because of difficulties completing one's education and maintaining employment.

Early onset of major depression or bipolar disorder may also impair upward mobility, but socioeconomic status also appears to affect the onset of major depression. A variety of studies suggest that the etiological significance of socioeconomic status in depression relates to the prevalence of major life stresses and persistent difficulties and the lower availability of coping resources and social supports¹⁵.

Disadvantages associated with socioeconomic status put persons with mental illness at risk for a number of negative outcomes that are not directly associated with the illness. Inadequate housing or homelessness remain significant problems for persons with mental illnesses and compromise efforts to provide meaningful services, yet we know very little about these issues in older populations¹⁶. Economic disadvantage is also a substantial barrier to access to health services and prescription medications in countries without universal health coverage¹⁷. These problems may be accentuated in later life for those with limited resources.

Socioeconomic status also helps to explain differences in mental health outcomes for different race and ethnic groups in the USA. While there do not appear to be major race/ethnic differences in the prevalence of severe mental illnesses independent of socioeconomic status, these factors significantly influence the course of illness. In the USA, rates of services utilization are lower for Mexican Americans, Asian Americans, and African Americans than Whites, when need for services is taken into account¹⁸. Economic barriers to access to health services contribute to this gap between need and service utilization among certain race and ethnic groups. Cultural differences in attitudes toward mental health services, perceived stigma and discrimination may also play a role¹⁹.

The importance of family for the onset and course of mental illness has been commonly examined. The onset of severe mental disorder early in life increases the likelihood of remaining unmarried, especially for men, and increases the risk of divorce and separation among those who do marry²⁰. Thus, many persons with severe mental illness will enter their later years without the benefits of a partner to provide emotional and instrumental support.

Styles of interaction within the family are also important. Most of the research has been focused on schizophrenia and suggests that a highly involved critical orientation to the patient contributes to relapse²¹. Instruction of families based on these principles has been found to be useful in some controlled trials, and offers a conceptual basis for psychoeducational approaches^{21,22}.

Increasing research in mental health focuses on stress and the coping process and the role of social support in either buffering the effects of stress or independently contributing to emotional well-being. While such factors as stressful events, coping resources, intimate and instrumental relations and self-efficacy have all been found to be associated with variations in psychological distress, their significance for major mental illness is less clear. Moreover, the role of these factors is condition-specific. In schizophrenia, stress acts as a trigger, affecting the occurrence and timing of episodes²³. In affective disorder, in contrast, stressful events and meanings assigned to these events appear to play a more causative role in conjunction with other factors¹². There is recent evidence that childhood adversity, such as separation from parents or sexual and physical abuse, may trigger onset of depressive disorders well into adulthood²⁴.

While, in general, it is believed that social support is important, the forms of social support useful to varying kinds of patients at

different stages of their conditions remain unclear²⁵. There is much research suggesting especially that intrusive forms of social interaction involving criticism may be detrimental to schizophrenic patients, and the relative protectiveness of intimacy, friendship and instrumental assistance in depression remain unclear. Social support remains a vague concept, having varying operational definitions. Contrasting views of social support explain some of the conflicting research results.

There is increasing research attention being given to the role of religion and religious participation in health outcomes for the elderly and a variety of studies suggest that different aspects of religious involvement play an important role²⁶. The pathways through which such effects are manifested remain unclear, although there is increasing suggestion that the social supports and instrumental assistance that might result from religious involvement contribute to better health. Research in this area confronts difficult selection biases, in that persons in better health may be more able and inclined toward religious participation and that persons drawn to varying religious practices may be different in their personalities, attitudes and behaviors than those with lesser or no participation. Nevertheless, the research in this area is gaining rigor and is suggestive of potential pathways.

The focus on coping skills as an important determinant of adaptation in more theoretically orientated studies has found expression in the development of psycho-educational programs to assist chronic mental patients in their rehabilitation. Although the research is modest, it supports the value of problem-orientated educational approaches that assist patients to manage everyday life situations better²⁷. Psychosocial research on the importance of self-efficacy has been translated into programs to increase patient empowerment. It has been suggested that such empowerment is an important feature of the success of some commonly studied rehabilitation programs²⁸. Increased empowerment may be an important factor, ameliorating the negative impact of highly regimented settings, such as nursing homes and other custodial institutions, but its therapeutic role in the normal range of life settings is yet to be established²⁹.

Social factors shape the processes through which individuals and families define illness, evaluate its meaning and significance and make decisions about needed care and appropriate practitioners. Epidemiological evidence shows that much serious mental illness is untreated, and those receiving treatment obtain services from a wide variety of practitioners. Processes of help-seeking are influenced by broad social beliefs about the nature of illness and what should be treated, characteristics of the individual and the social contexts in which mental illness occurs, and the organization of services and their physical, social and economic accessibility¹. Members of varying age cohorts have been socialized differently in relation to the recognition of symptoms, appropriate sources of help and the social stigma of seeking care for particular types of problems. Selection from the community to varying types of service providers is a two-stage process, which depends on general factors affecting the inclination to seek assistance, and other factors more specific to the choice among alternative practitioners. A large proportion of all patients receiving care for a mental illness receive such care exclusively from general medical practitioners, and decisions affecting referral to the specialty mental health sector result not only from personal definitions and inclinations of patients and their families, but also from the organized pathways within a health care system, the ability of generalists to recognize mental illness in their patients and their attitudes to specialty mental health services. Patients with mental illness are referred more readily to specialized services when illness and illness behavior imply social risk and disruption³⁰. Diagnosis, itself, is a poor predictor of the referral process.

Patterns of help seeking differ by age. The elderly are less likely to seek psychiatric care than younger adults, and probably are

more reluctant to report affective symptoms to interviewers or physicians. In contrast, the elderly complain commonly in general medical settings of diffuse physical symptoms and vegetative symptoms characteristic of depression, and have relatively high rates of receiving prescribed psychoactive drugs. While somatization, as measured more formally, does not seem to vary by age, the elderly are more likely to present distress in a somatic idiom. Interpretation is complicated by the fact that the elderly have higher rates of ill-health and chronic disease than younger individuals, and it is difficult to sort out physical concomitants of chronic disease from somatization of psychosocial distress and depression. Psychoactive drug use also reflects physician behavior, which may be shaped by stereotypes of the elderly and other factors.

The social response to mental illness is influenced by such factors as general attitudes, values and ideologies, concepts of the nature of disorders and their causes, available treatment technologies, the structure of health and welfare services, and the system of social entitlements that a society makes available. The deinstitutionalization of the mentally ill has followed a different course in varying countries, but each of the above factors plays some role in every instance. In the USA, large-scale deinstitutionalization only became possible in the middle 1960s with the expansion of welfare programs that provided subsistence and payment for alternative residential care for mentally ill patients in the community. Some of the evident difficulties in community mental health services reflect cutbacks or fluctuations in welfare and housing entitlements¹.

The magnitude of serious mental illness depends on the distribution of the population, and the numbers of persons in age groups at risk for varying diseases and disabilities. The burden of mental illness depends not only on its magnitude but also on the types of social institutions and programs that help insulate patients, families and communities from its most disruptive stresses. The elderly population, and persons at very old ages, are growing in the USA and many Western countries. The dementias are increasingly important, and patients with such disabilities constitute a growing proportion of the severe mentally ill population. Depending on the constellation of institutional services and home-care programs, patients with dementia are treated in a variety of settings, some placing very large burdens on families and friends. The distribution of such burdens, and the definition of responsibilities, is a political process and a key social policy issue throughout the world.

For the last several decades, the nursing home has been the setting for care of persons who are greatly restricted in the activities of daily living, and a large proportion of persons in these settings have dementia and depression as a secondary condition³¹. Typically, admission to nursing homes occur when individuals are incapable of caring for themselves, when their physical and psychiatric problems create unmanageable burdens for their caretakers, or when community caretakers are no longer available. Admission is often triggered by such events as significant loss of function following trauma, such as hip fractures, confusion and wandering, and incontinence. Relatively few elderly with a primary diagnosis of mental illness, without dementia, are in nursing homes, although nursing home admission often exacerbates confusion, apathy and depression. Most elderly persons resist nursing home admission as long as they can, and increasingly alternative community settings and home-care services are provided to prevent or delay such admission³².

Nursing homes, like the traditional long-term mental hospital, contribute to an institutional syndrome resulting from vulnerabilities of patients as they respond to decreased social participation, sensory deprivation, loss of efficacy and control over daily life decisions, institutional routinization and the like. There is persuasive evidence that efforts to keep patients involved

and participating in valued activities help maintain both mental and physical function. Many nursing homes fail to give these social aspects adequate attention, and patients spend much of their time in isolation and uninvolved. Nursing care may be the best treatment context for some patients who are seriously incapacitated and require a broad range of services difficult to provide in the home or in other community settings, but most impaired elderly benefit from settings that are more successful in sustaining independence and social engagement.

The capacity to cope effectively with mental illness depends on the organization of kinship groups and households, and the existing social institutions in a society. The growing prevalence of divorce, single-person households, small families, high female participation in the workforce, geographic mobility and other trends make it difficult to put increasing reliance on informal social networks for caretaking. Developing alternative structures that are financially viable and humane constitutes a growing challenge everywhere.

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The Natural History of Psychiatric Disorders: Early-onset Disease in Late Life and Late-onset Illness

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Is psychiatric pathology fundamentally different depending on when in the life course it emerges? Is the heterogeneity that characterizes late life psychiatric disorders—their risk factors, presentation and course—partly resolved by distinguishing between ages of first onset? This chapter summarizes why studying the onset of psychiatric illness is so difficult and, in spite of that difficulty, what distinguishes the epidemiology of early-onset disease in late life from late-onset illness.

With respect to age of onset, the best-characterized DSM-IV disorder (and our primary focus in this chapter) is major depressive disorder (MDD). Some attention will also be directed to bipolar (BP) disorder and schizophrenia. Much less research has distinguished between early and late onset of dysthymia, “late paraphrenia”, or delusional disorder; we touch only briefly on these subtypes. We do not discuss depressive and/or psychotic prodromes of dementia or psychiatric symptoms of other organic etiology, such as cancer or multiple sclerosis, with the exception of disorders related to cerebrovascular infarcts and depression secondary to cardiovascular disease.

Studies of the prevalence and correlates of early-onset vs. late-onset disorders are treacherous for methodological reasons that plague all psychiatric studies and all studies with retrospective designs: determining *who* has *what* disorder and *when* it was first detected. Sometimes there is no consensus on the criteria for a “case”, because psychiatric disorders are based on clinical signs and symptoms and represent heterogeneous underlying pathophysiological and psychosocial causes. Subjects may be considered a case according to one set of criteria but not according to another. Elderly persons in particular report significant psychiatric symptoms without meeting diagnostic criteria. Not infrequently, emergent symptoms may alter a diagnosis of long standing. Thus, clinical populations, from which many study samples are drawn, may be non-representative of all affected elders, due to varying patterns of help-seeking, clinical referral and symptom trajectories. All elderly populations in communities and institutions, are subject to survival of the fittest, and early-onset subjects may be systematically absent due to death.

Determining the precise age of onset is also problematical. Recall of specific dates may be compromised by memory lapse or telescoping of events, cognitive impairment or a “reminder effect” associated with multiple episodes. Classification of early-onset vs. late-onset depends on the cut-point chosen, which is often unspecified in diagnostic schema; across various studies, cut-points have ranged between 20 and 60 years of age, although most

use 45 or 50 years. For all of these reasons—diagnostic variability, selection bias, subject attrition and classification issues—estimates of the prevalence and risk of early-onset and late-onset disorders vary dramatically, as described below. Nevertheless, this chapter describes the state of agreement regarding differences in early and late affective and psychotic disorders.

AFFECTIVE DISORDERS

Major Depressive Disorder

The prevalence of affective disorders (MDD/BP) is lower among community-dwelling elders than among younger populations^{1,2}. Prevalence estimates of late-life MDD from large studies (*n*51000) range between 1% (USA) to 3.7% (Finland), and prevalence of minor depression ranges from 8.3–23.2%^{3–5}. In nursing homes, 5.9–16.5% of patients demonstrate clinically significant depression^{6–7}. Among medical outpatients and inpatients, respectively, 5–7% and 10–20% may meet criteria for MDD^{8–10}. Of affected elders in the community, van Ojen and colleagues¹¹ estimated that 70% may have had a first onset after the age of 65, i.e. late-onset depression (LOD) (Amsterdam, Holland). In the ECA (USA) study, the mean age of onset of MDD for subjects older than 65 years was 48.9 years, suggesting a large proportion of patients with late-onset¹.

The most provocative etiological findings regarding age of onset concern structural and functional brain abnormalities among LOD patients. Depressed elderly patients tend to have more abnormalities, particularly in the frontal regions, than do controls, although not universally^{12–14}. LOD patients tend to present with more brain lesions than do patients with early onset of depression (EOD)¹², and subcortical and basal ganglia changes have been associated with greater severity in LOD¹⁵. These results from MRI studies are suggestive when paired with results showing increased vascular medical co-morbidity (e.g. hypertension, stroke and heart disease) among LOD patients^{13,16}. Thus, risk of some subtypes of LOD may be attributable to vascular impairment and, by implication, preventable, using heart-healthy strategies such as diet, exercise and stress management. Where these results are heterogeneous, the inclusion or exclusion of subjects with cardiovascular histories may be a major factor^{16,17}.

At the other end of the modifiability spectrum, genotype may account for differences in the causal webs of EOD and LOD.

Krishnan and colleagues¹⁸ found patients with the *ApoE3/4* allele to have later age of onset than those with *2/3* and *3/3*; conversely, Holmes *et al.*¹⁹ showed that *ApoE2* protected against early onset, thus delaying the onset of depression into later life; other studies were inconclusive²⁰. Although these findings conflict, the potential to locate common genetic risk factors for depression and dementia in late life continues to receive empirical attention. Pedigree studies have been much more univocal: elders with EOD consistently report more affected close relatives than do elders with LOD^{16,21}. Some have suggested recently that age of onset may be a proxy for multiple risk factors, including structural brain changes, vascular medical co-morbidity, neuropsychological impairment, and family history of mood disorders²².

Evidence of other non-medical risk factors for EOD and LOD is preliminary. Single studies show males and African-Americans at greater risk of LOD than females and Whites, respectively^{23,24}.

The clinical presentations of early-onset and late-onset disease among elderly patients are not strikingly different, with the possible exceptions of more apathy, anxiety, and psychotic features in LOD²¹⁻²⁵. Most neurological tests also show no differences, with the possible exception of more impaired executive memory and visual naming ability in LOD^{26,27}.

On balance, prognosis may be slightly worse for EOD elders, although there are serious methodological difficulties with the evidence and significant heterogeneity of course in late life. Cole²⁸ summarized conflicting early findings of differential disease trajectories, and studies over the subsequent decade have continued to conflict, some showing EOD elders to have a more relapsing course, more frequent episodes and more treatment refractoriness, and others showing LOD elders to have less favorable treatment response, more cycling in and out of symptoms, and more psychiatric co-morbidity²⁹⁻³¹. Still other studies demonstrate no difference. However, when treatment and outcome criteria have been standardized, neither LOD nor EOD elders appeared at increased advantage relative to absolute rates of 1 year outcomes (remission, relapse, recovery or recurrence), although weeks to remission were greater in one sample of LOD elders²².

In a recent comprehensive review of the mortality of depression³², only one of 57 studies compared mortality by age of onset. Rabins *et al.*³³ found no differences in mortality by age of onset in a small study. More recently, Philibert *et al.*³⁴ reported lower survival rates for subjects with LOD, especially women with onset later than their 6th decade.

Bipolar Disorder

Manic symptoms and bipolar disorder (BP) I and II are rarer in the population than depressive symptoms or MDD and are most rare late in the life course³⁵. Fewer than 1% of US elders in the ECA study reported a 1 week period or more of elevated, expansive or irritable mood; lifetime prevalences of these disorders among persons 65 years and older were 0.1% for BPI and PBII in the same study¹. Evidence from the UK suggests that the incidence of mania may be bimodal, with a detectable peak after 60 years^{36,37}. Among elderly hospitalized psychiatric populations, 3.4–6.4% of all admissions and 12.5% of admissions for affective disorders present with mania; 12–16% of general hospital patients qualify for a bipolar diagnosis³⁵.

Historically, the two potential causes of mania were thought to be stressful life events and organic disease. Kraepelin³⁸ first noted the precipitating role of stressful life events in manic depression, and subsequent studies have implicated such events in the incidence of bipolar disease, particularly early-onset mania (EOM)³⁹. Late-onset mania (LOM), on the other hand, has long been associated with both vascular and cerebral anomalies and their correlates, including obesity, head injury, EEG changes and

dementia, with confirmatory evidence in more recent studies comparing older vs. younger bipolar patients⁴⁰ and LOM vs. EOM patients^{39,41}. Yassa *et al.*³⁵ suggests that where significant life events distinguish LOM patients from aged controls, an organic substrate may have predisposed affected elders to a vulnerability to such events, and they then “react with a ‘catastrophic’ response (mania)” (p. 126).

A positive family history of bipolar illness has been more characteristically linked with EOM than with LOM patients^{39,42-44}. However, methodological problems render this generalization suspect. Retrospective designs, low age-of-onset cut-off points and failure to distinguish between onset of depression vs. onset of mania are a few of the problems limiting the evidence⁴³. Evidence of a gender differential in risk of early-onset or late-onset bipolar disorder has also been inconclusive³⁹. Interracial comparisons of age of mania onset are not available.

Findings of differential clinical presentations of LOM and EOM in late life have not proved robust. The stereotype that LOM presented more often with “slow and fragmented flights of ideas, more rarely with elated affect, but more often with paranoid and aggressive features (p. 120)” was not confirmed in a summary of studies comparing the phenomenology of LOM and EOM in late life³⁵.

Prognostic differences by age of first onset of mania are difficult to substantiate⁴⁵⁻⁴⁷. In particular, differential survival may underlie the evidence of poor outcomes associated with LOM. To the degree that healthy social interactions are relevant to managing the course of bipolar disorder, recent preliminary findings are noteworthy: age of mania onset was higher among subjects with more positive assessments of social support and receipt of informal instrumental support^{39,45}.

PSYCHOTIC DISORDERS

Like affective disorders, late life psychoses (LLP) are heterogeneous, and the reach for consensus on diagnostic terminology has been torturous. In this summary we primarily discuss early-onset vs. late-onset schizophrenia (EOS/LOS), with brief mention of delusional disorder and entirely omitting discussion of alternative nomenclature (e.g. late paraphrenia) or subtypes less well characterized with respect to age of onset (e.g. schizo-affective and other psychotic disorders).

The lifetime history of schizophrenia is probably 41% in the elderly community population⁴⁸, with annual incidence of three new cases (and 45 relapses) per 100 000 elders⁴⁹. Approximately 25% of elderly schizophrenics experienced a first episode after age 50⁵⁰. Onset of delusional disorder, on the other hand, occurs later, with average onset in the 40s among men and the 60s among women⁵¹. Late-life incidence of delusional disorder is 15.6/100 000; prevalence in the elderly population is 0.04%⁴⁹.

Two powerful risk factors for LOS appear to be cerebral atrophy and brain injury, although a significant number of patients with late-onset LLP present with no apparent cerebral pathology^{52,53}. Compared to affective disorders, there is less evidence that EOS and LOS elders present differently with respect to MRI findings⁵³ or cardiovascular medical co-morbidity. A positive family history of psychosis appears incrementally more likely among, respectively, non-psychotic elderly controls, elders with LOS, and elders with EOS⁵⁰. Findings concerning ApoE differences in onset or phenotype are limited and inconclusive⁵⁴. Female gender and sensory impairment may predispose to LLP^{48,55}.

There is considerable heterogeneity in the outcome of schizophrenic episodes. A significant number of schizophrenic patients improve and recover fully; others relapse intermittently; others remain chronically symptomatic^{56,57}. Long-term studies include

mostly EOS patients, who make up the majority of geriatric schizophrenics. LOS, on the other hand, portends generally poor outcomes, although social adaptation may be better in this group, compared to EOS^{50,58,59}.

CONCLUSION

Overall, there is growing evidence that older patients with late-onset psychiatric disease are different from those who come to late life with a history of early-onset psychiatric illness. Their varying co-morbidity and symptom profiles suggest different (although potentially interacting) etiological factors, with genetic and/or social environmental factors predominating in early-onset disease and behavioral factors in later onset. More work is needed to describe any differential course, but social adaptation appears better for elders whose disease experience has been shorter.

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Mortality and Mental Disorders

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INTRODUCTION

The topic of mortality and mental illness has a long history. Even if we discount the pioneering nineteenth century study of Farr¹ we can still go back as far as the study of Ødegård² among the contemporary studies. Both of these were, of course, patient-based and covered all ages.

Why should anyone want to study mortality and its relationship to psychiatric illness in later life? There are many questions we might want to ask:

- Are older people with psychiatric illnesses more likely to die than those without?
- Are older people with psychiatric illnesses more likely to die than younger sufferers after allowing for the general rise in the force of mortality with age?
- If they are more likely to die, in either case, why is this?
- Whether they are more likely to die or not, do they die from different causes?
- What theoretical issues does their risk of mortality raise?
- What implication for service provision does their risk of mortality, elevated or not, have?

There have been some attempts at studying these issues and a small number of relevant reviews. In this chapter there is no attempt to provide a bibliography of all the empirical studies. Instead a number of key references are chosen to illustrate themes.

DEATH CERTIFICATE STUDIES OF MORTALITY

A number of studies have used death certificate data to study mortality and the dementias. The primary goal of these studies has been to estimate prevalence without the expense of performing a community study. Only one study, that of Flaten³ in Norway, gave estimates of prevalence close to those from community studies, and it has been generally accepted that such studies tell us about the death certification habits of doctors but rather little about the prevalence of dementias or the relationship between the dementias and mortality. There have been no such studies for illnesses other than the dementias.

PATIENT-BASED STUDIES

A full review of these was undertaken by van Dijk and colleagues⁴ which should be referred to for details of this work. For reasons outlined in the next section, these studies will not be further discussed here.

COMMUNITY-BASED STUDIES

The primary source of information about the link between mortality and mental illness in this age group is the increasing number of community studies. Careful follow-up of well-defined series of carefully diagnosed cases has its place, but the selection that goes on between primary and secondary care gives rise to the risk that the information cannot be generalized. Since most cases of dementia and depression are managed in primary care, it is there that we must seek information on the mortality of cases of mental disorder. One of the most comprehensive reviews of the past few years was that of Schröppel⁵. She concluded that relatively little was known for either disorder.

The years since Schröppel's review have seen an explosion in studies relevant to dementia and depression. Two recent systematic overviews have synthesized those community mortality studies for dementia⁶ and depression⁷. Disappointingly few of the primary studies provide effect sizes that can be combined, but there are enough to warrant the exercise for both disorders. For dementia, meta-analysis of six studies gave an odds ratio for mortality of 2.63 (95% confidence interval 2.17–3.21). There was weak evidence of a higher risk for vascular compared to Alzheimer's, for increasing risk with increasing severity of dementia, and for decreasing relative risk with age. There seemed no evidence of a sex difference. For depression, meta-analysis of 15 cohorts gave an odds ratio of 1.73 (95% confidence interval 1.53–1.95). There was weak evidence that men had a higher relative risk and that studies with longer follow-up have a lower risk.

There have been few studies of other diagnoses in older age, and there has been no systematic review of them. An account is given by Langley in her review⁸.

CAUSE OF DEATH

Only one study⁹ appears to have addressed the problem of the cause of death in community studies of dementia. Prior to this there had been a number of studies of patient cohorts. Although these doubtless have value, they do not help to solve the problem of the causes of differential mortality. For that we require a direct comparison between the proportion who die of cause X in the demented group with the proportion who die of cause X in the non-demented group. It seems so obvious that this is needed that it is surprising that only Jagger and her colleagues⁹ have addressed the issue. They found that, relative to the general population, Alzheimer's disease sufferers had a lower risk due to age, for being manual social class, and for having a history of cancer, and a

higher risk for being male. This concurs with the meta-analysis quoted above for age, but not for sex.

A detailed editorial¹⁰ discusses issues in research into depression and mortality. In particular there is difficulty in separating association from causation.

METHODOLOGICAL ISSUES

A number of measurement issues have failed to receive appropriate attention. The studies of community samples use different methods of measuring differences in risk. Although for low risks the numerical difference between risk ratios and odds ratios is small, comparison would be simpler if studies reported in a common way.

Patient studies have not really addressed the issue of bias introduced by differential identification of the patient group and the controls. For instance, if date of diagnosis is used as the starting point for survival analysis, there will be a survivor bias and it will be difficult to identify controls in a similar way.

Studies comparing patient groups with the general population have also failed to account for the fact that demented people are part of the population, and if their death rate is very different from that of the population as a whole, and if they form a substantial part of the population, as they will beyond age 85, then the death rate quoted for the general population is an overestimate of that expected for non-demented people.

Few studies have put their results into any clear framework. For instance, the possibilities raised by epidemiologists¹¹ studying chronic physical illness, and implicit in the textbook relationship between prevalence, incidence and duration of illness, have hardly been explored¹².

DISCUSSION

Mortality has been one of the most studied endpoints in epidemiology, but the reader of the community studies will be struck by the fact that, for few of them, mortality was a primary end point. This has undoubtedly created the patchwork of methods of reporting which bedevils synthesis.

There has been a considerable literature relating mental disorders in younger adults to mortality¹³, which has confirmed an increased risk for affective disorders (typical SMR 1.4) and

schizophrenias (typical SMR 2.3). The studies quoted here suggest a similar, possibly slightly greater, risk for depression.

There is clearly an elevated risk for both dementia and depression, but there is little that can be confidently asserted beyond that. We do not know whether the risk is modified by other possible explanatory variables like age, sex, physical illness of functional status, neither do we know what causes of death account for the excess cases.

As yet, we do not know what would happen to mortality if we treated depression in older age more vigorously, or what would happen if universally effective treatments for dementia became available and we could treat people with dementia.

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Long-term Outcome Studies of Psychiatric Disorders: Methodological Issues and Practical Approaches to Follow-up

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What are the long-term outcomes of various types of psychiatric disorders? How do individuals, interacting with their environments, shape the course and consequences of mental illness? What environmental, personal and illness characteristics are associated with recovery and improvement? Are such characteristics disorder- or life stage-specific? A growing number of long-term follow-up studies have begun to address such questions. Their results both confirm and challenge accepted wisdom. Long-term outcomes of schizophrenia, for example, do seem to be worse on average than those of affective disorders¹; at the same time, however, outcomes of schizophrenia are remarkably varied, undermining conceptions of its course as progressively deteriorating and chronic². Substantive results from long-term follow-up studies are included in other chapters of this volume (*see specific disorders*). This chapter focuses instead on several basic methodological issues—case identification, co-morbidity, choice of comparison groups, sample representativeness and measurement issues—as they pertain to long-term follow-up studies (*see refs^{3–6} for elaboration*). Our purpose is to alert readers to issues that potentially influence the interpretation and comparison of results. Issues associated with the statistical analysis of longitudinal data are beyond the scope of this chapter and are not included (*see refs^{6–8}*).

There are two basic types of follow-up investigations: (a) prospective studies, in which the investigator defines a sample on the basis of current attributes (e.g. psychiatric or exposure status) and follows the sample forward in time; and (b) retrospective cohort designs, in which the researchers define the sample on the basis of some past characteristic (e.g. hospital admission during a specified time period, participation in an earlier study) and then reconstruct their subsequent life course up to some later point in time, using records, retrospective interviews, etc.³ Retrospective cohort (or “catch up”) designs require less time to complete than prospective studies covering a similar duration, but are more limited by the availability and quality of extant data and problems of recall. Both types of investigations have been used to elucidate the long-term course⁹ and outcomes of psychiatric disorders¹. The issues discussed below apply to varying degrees to both types of investigation.

CASE IDENTIFICATION

Case identification refers to the criteria and procedures used to determine what constitutes a case of the disorder under study and

addresses the question, “long-term outcomes of what?”¹⁰. Without explicit specification of what constitutes a case, similarities and difference in observed outcomes across studies are difficult to interpret. For example, observed differences may reflect discrepancies in diagnostic practice; observed similarities may be illusory. Case identification is particularly problematic in long-term follow-up studies, where baseline data and diagnoses often predate the introduction of modern diagnostic systems and standardized assessment tools. One way in which investigators have dealt with this problem is to rescore baseline data (e.g. from hospital charts, case notes or interviews) using one or more current diagnostic systems (e.g. DSM-III-R, ICD-10) and to select and compare cases based on these newer diagnoses⁴. This practice allows for comparison of outcomes using different nosologies and, insofar as criteria for inclusion and exclusion are reported, facilitates comparison of results across studies. Such benefits of rescoring, however, remain dependent on the quality and comparability of baseline data. Signs and symptoms that were never attended to or recorded obviously cannot be recovered, and their absence potentially introduces error into the rescoring process.

CO-MORBIDITY

A second issue concerns co-morbidity (i.e. the co-occurrence of two or more psychiatric disorders in an individual) and addresses the question, “to what extent are observed outcomes due to psychiatric conditions other than the one under study?”^{5,22–25}. What appear to be differences in outcomes of schizophrenia, for example, may be due in part to differences in the prevalence and combination of secondary conditions. Here, too, the assessment of co-morbidity and its ramifications is particularly difficult in long-term follow-up studies, because outcomes may be influenced not only by secondary disorders recorded at baseline but also by the onset and recurrence of other disorders between baseline and follow-up. While some assessment tools [e.g. Schedule for Affective Disorders and Schizophrenia—Lifetime (SADS-L)] generate the data needed to make lifetime diagnoses, sample attrition and reliance on informants may impede reliable assessment of interim conditions for substantial numbers of subjects.

CHOICE OF COMPARISON GROUPS

The counterpart of case identification is the designation of comparison groups and addresses the question, "long-term outcomes of disorder X compared with what?" Comparison groups provide benchmarks for interpreting measures of outcome. What comparison groups are appropriate depends, of course, on the research question under investigation and includes, among others, samples of "normal" individuals, of individuals with other psychiatric or medical conditions, or population norms, if available. Comparisons may also be made on the basis of some risk factor (e.g. age of onset) or treatment of interest. What is important to remember when selecting comparison groups and interpreting results is that members of the case, or target, group may differ from those of the comparison group in ways other than the disorder or risk factor of interest, and these "other ways" may account for subsequent differences in outcome and course. Thus, it is important to assess whether comparison groups are comparable to case groups on extraneous dimensions that could influence outcomes.

SAMPLE REPRESENTATIVENESS AND ATTRITION

The representativeness of the sample at baseline and follow-up addresses the question, "to what extent and to whom are results generalizable?" Most current long-term follow-up studies are based on patient samples selected from hospitals, other treatment settings, clinician caseloads or case registries. Selection into treatment, however, is not a random process¹¹. Treatment rates have been shown to vary by type of disorder¹², social and demographic characteristics of individuals^{12,26}, historical period¹³, service delivery system and country. In addition, within a catchment area, patients are sorted and filtered into different types of treatment settings (e.g. public vs. private). Thus, the life trajectories of treated subjects may differ not only from those of untreated cases but also from those of patients treated in dissimilar settings.

More significant is the problem of sample attrition, whether due to death, inability to participate, refusal or failure to trace. This is because subjects lost to follow-up tend to differ in systematic ways from those who are located and interviewed^{3,14}. Researchers often attempt to assess the type and degree of bias by comparing respondents and non-respondents on baseline characteristics; however, as Kelsey *et al.*³ note, "the only way to ensure that bias stemming from loss to follow-up does not distort the study results is to minimize losses through intensive efforts to locate each cohort member" (p. 109). In addition, by reducing sample size, attrition reduces the statistical power of analyses to reliably detect outcome differences. With effort, several long-term follow-up studies have successfully minimized sample attrition¹⁵; the training and organizational strategies used have been summarized^{15,16}.

MEASUREMENT OF OUTCOMES

The measurement of outcomes is clearly central to long-term follow-up studies and raises the question, "on what dimensions and how should outcomes be assessed?" Many different measures of outcome have been used in follow-up studies, including measures of symptomatology, hospitalization, role functioning, impairment of social relationships and recovery^{1,17,18}. Reliance on single global indices of recovery have proved largely unsatisfactory. Such indices are not only difficult to compare across studies but also imply an overly uniform picture of subjects' functioning. Subjects who fare poorly in one life domain (e.g. employment)

often fare better in others (e.g. symptom profile)¹⁸, and such discrepancies are masked by single global indicators of recovery. Even seemingly objective, domain-specific measures (e.g. unemployment or rehospitalization rates) can be difficult to compare across studies if they are influenced by setting specific economic conditions or social policies^{4,19}.

Sample attrition further complicates the measurement of outcomes. How should outcomes be assessed for subjects who are deceased or unlocated at follow-up? Investigators often rely on informants and/or records to provide information. However, in addition to raising ethical considerations, informants not only differ from subjects in their access to (and probably recall of) relevant information but also assess subjects' status from different perspectives. Records, too, may not contain the data of interest and, like informant reports, differ in perspective from subjects' self-reports^{3,26}. In light of such considerations, it is important to follow the advice of Bromet *et al.*⁴ and ". . . be cautious in comparing results across studies without carefully ascertaining both the definition of outcome and the criteria used in implementing that definition" (p. 154) (*see refs*^{3,17} for elaboration).

MEASURES OF INTERVENING VARIABLES

The longer the time interval between baseline and follow-up, the more likely that intervening events and circumstances play a role in determining outcome status. Measurement of such events and circumstances addresses the question, "what are the interim processes and mechanisms that account for observed outcomes?" Here prospective designs have an advantage over retrospective cohort studies because measurement of potentially mediating and confounding variables can be explicitly built into both baseline and follow-up data collection, and time intervals between follow-ups can be chosen with intervening mechanisms in mind. Retrospective cohort designs are more dependent on the availability of records and/or the recollections of subjects and informants, which are vulnerable to recall and reporting biases^{3,20,21,23}. Problems associated with reporting and recall are not avoided by prospective studies, however, especially when intervals between follow-ups are long.

PRACTICAL APPROACHES TO FOLLOW-UP

Smith and Watts²⁷ provide a thorough summary of methods for locating absent and deceased subjects. Their review of available procedures emphasizes the use of electronic databases as effective means for identifying the location and vital status of lost subjects. Various tools for locating participants lost to follow-up are currently available.

Sources for Locating Patients Assumed Alive

The US Postal Service is a convenient source of information when investigators are faced with outdated addresses. Upon request, an updated address can be obtained as long as the request is made within the year that the individual moved.

Incorrect telephone numbers can be updated by contacting the telephone operator for information on new telephone listings. Unlisted telephone numbers may be obtained, depending on the policy of the telephone company. White-page listings for the entire USA are available for purchase on compact disk (CD-ROM). Compact disks are updated yearly and are relatively affordable. Telephone numbers can also be updated by use of the

crisscross (or reverse) directory, which lists telephone numbers by address.

The Internet serves as an effective and cost efficient method of locating absent subjects. Various websites contain an extensive amount of information regarding residential listings. Internet sources differ by the type of information required to perform a search. When utilizing online databases, Smith and Watts²⁷ suggest that: "It is important that the investigator try each source before moving on to the next site. Failure to locate a number on one site does not mean that a search of another site will be fruitless" (p. 434). Some examples of current websites include:

www.infoseek.com
 www.semaphorecorp.com
 www.databaseamerica.com
 www.four11.com
 www.yahoo.com/reference/whitepages
 www.angelfire.com
 www.yahoo.com/search/people
 www.switchboard.com
 www.whowhere.com
 www.lookupusa.com

The Department of Motor Vehicles can provide useful information on an individual, including one's address, date of birth and social security number, and can select physical characteristics; however, investigators must be cognizant of state-specific limitations and restrictions on public access to information. Similarly, voter registration files are restricted in certain states, yet careful review of files, when available, can produce useful information.

TRW, TRANSUNION and EQUIFAX, the nation's major credit bureaux, can provide vital information on patients, such as current and previous addresses, social security number, birth date, employer and spouse's name. Each of these credit-reporting services offers affordable products for assessing patient status. TRW offers TRW Social Search and TRW Address Update. Transunion offers TRACE, TRACE^{plus}, ReTRACE and the ATLAS. Equifax offers DTEC and ID REPORT. Each product can successfully provide essential data but differs by the specific information generated and what is required for initiation of a search. Utilization of these credit bureaux is legal, offering comprehensive, affordable, time- and cost-efficient means for locating patients.

Information brokers may also be of use in attempting to locate missing subjects. Information brokers have access to varied sources of information, yet are typically costly options. As a result, it is recommended that researchers assess the necessity to locate their subjects prior to selecting such alternatives^{27,29}. Similar sources of information may be available outside the USA.

Sources for Locating Patients Assumed Deceased

Various services are available for locating individuals who are believed to be dead. The National Death Index (NDI) identifies the states in which death occurred, the corresponding death certificate number and the date of death. The NDI file provides information for all 50 states, the District of Columbia, Puerto Rico and the Virgin Islands. The NDI database is updated annually and begins with deaths occurring in 1979.

NDI users can apply for NDI^{plus}, an optional service that, in addition to the aforementioned information, provides details on the underlying and multiple causes of death for deceased patients. Procedures for requesting information through NDI vary by state. Prospective NDI and NDI^{plus} users must submit an application form to the National Center for Health Statistics. Information regarding NDI and NDI^{plus} can be obtained by contacting:

National Center for Health Statistics, Centers for Disease Control and Prevention, 6525 Belcrest Road, Room 820, Hyattsville, MD 20782-2003, USA.

Tel: (301) 436-8951, ext 109 or 101; fax: (301) 436-7066; e-mail: ROB3@CDC.GOV

The Equifax Nationwide Death Search provides information on deaths since 1955 from data compiled from the Social Security Administration. Search results will produce birthdate, date of death and state and zip code at death. Equifax is updated continuously and is therefore recommended for obtaining data regarding recent deaths. Furthermore, because records are available beginning in 1955, Equifax is potentially more useful for some retrospective cohort designs than the NDI²⁸.

Other sources for identifying deceased subjects include the Death Master File from the Social Security Index^{27,29}. This information can be accessed at www.ancestry.com.

Contact information for additional services includes:

Social Security Administration's Death Master File, US Department of Commerce, Technology Administration, 5285 Port Royal Road, Springfield, VA 22161, USA.

Tel: (703) 487-4630

Equifax Credit Information Services, PO Box 105835, Atlanta, GA 30348-5835, USA.

Tel: (800) 944-6000

Trans Union Corporation, PO Box 8309 File 99506, Philadelphia, PA 19101-8309, USA.

Tel: (610) 690-3126

CONCLUSIONS

Long-term follow-up studies provide a useful vehicle for investigating stability and change in the course and consequences of psychiatric disorders; they raise a number of methodological and practical considerations as well. In this chapter we have tried to highlight basic methodological issues and practical approaches that bear on the interpretation and comparison of research findings and to reference practice approach and texts that address the issues in greater detail.

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Part D

Diagnosis and Assessment

- DI Epidemiology, Diagnosis and Nosology
- DII Clinical Assessment
- DIII Standardized Methods and Rating Scales

The Importance of Multidimensional Assessment in Clinical Practice

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In many ways, geriatric psychiatry is a paradigm for contemporary medicine. A multidisciplinary team is required to address the complex range of assessment and treatment issues. After all, older people usually have more than one medical illness, are on several medications and may have considerable limitations (vision, hearing, mobility, cognition, finances).

According to Eastwood and Corbin¹, “the problems presented in the assessment and treatment of geriatric psychiatry patients may be best solved by a team composed of members with diverse professional training, meeting frequently to plan investigations, discuss findings and formulate coherent and comprehensive treatment plans. Such a group must function with little overlap of tasks and few interdisciplinary jealousies, be problem-orientated and exercise communication skills in order to be effective and efficient. Although the physician typically chairs team meetings and takes medical and legal responsibility, each member of the team must make a definite contribution, either in direct patient care, or with technical and back-up services”, and “Whatever the level of sophistication, a geriatric psychiatry unit should be able to provide an holistic and eclectic approach to assessment of the patient”.

For each patient there may be several assessments, each with a definite purpose. Pre-admission assessment in the home is valuable for evaluating functional level and the appropriate level of care. Admission assessment may be necessary to make a diagnosis and treatment plan. Follow-up assessment helps monitored treatment effects and overall outcome. The following health care professionals are required in the therapeutic team:

1. *Psychiatry.* Geriatric psychiatry is best practiced in a short-stay, 10–15 bed unit. Ideally, the unit should be free-standing for the elderly alone. Some would agree, because of the medical risks and need for specialist consultations, that the unit should be in the general hospital. The psychiatrist brings together the psychosocial and biological aspects of medicine and is preferably a geriatric psychiatrist. The amount of training received in a general psychiatry residency is inadequate to qualify as a geriatric psychiatrist. Conversely, a full fellowship is not necessary and a separate 6 month rotation is probably sufficient.
2. *Primary care physician.* It is useful to have a primary care physician on the team. This person can help identify medical problems, and suggest tests and medication to deal with these. They can help in the initial phases of assessment. The pace of research makes it difficult for a psychiatrist to keep up satisfactorily with new medications in other specialties and even for common illnesses like hypertension and diabetes.

3. *Consultation team.* It is useful for the geriatric psychiatry group to offer consultation services to other parts of the hospital. Looking for cases of delirium, dementia and depression to treat or to transfer to the geriatric psychiatry ward is a useful way of doing consultations and sharing good relations with other specialties.
4. *Nursing.* Optimally, the same nurse should be responsible for initial contact, inpatient care and follow-up. With this familiarity, the nurse can be responsive to presenting problems of safety and security, prostheses, fluctuations in daily activity, signs of therapeutic response and drug or other sensitivities.
5. *Social work.* The social worker can assess the strengths and weaknesses of the patient's current social network, and evaluate the potential for benefit from individual, family and group psychotherapy. The social worker should be able to provide individual support to the patient and family members.
6. *Occupational therapy.* The occupational therapist can evaluate the patient's response to social, physical and intellectual stimulation and self-care and instrumental activities of daily living. Occupational and leisure needs can be linked to community agencies upon discharge. This therapist should be able to help patients and their families with activities (music, storytelling, group activities, such as playing various games) that can be used to help cope with problem behaviors and emotional distress.
7. *Psychology.* The psychologist can critically determine global and specific intellectual deficits and help manage certain behaviors.
8. *Pharmacy.* The pharmacist is vital in monitoring drug levels, interactions and reactions in order to avoid the problem of delirium.

It is also advisable to have the consulting services of such specialties as internal medicine, cardiology, neurology, ophthalmology, otolaryngology, urology, physiotherapy and dietary science.

The team faces a variety of diagnostic syndromes. These include dementia, affective disorder, delusional disorder, neurosis and personality disorder, alcohol or drug abuse, and delirium. Each of these requires a standardized diagnostic, treatment and management protocol, all subject to measures of efficiency and efficacy.

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Classification of Dementia and Other Organic Conditions in ICD-10

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The classification of mental disorders in the Tenth Revision of the International Classification of Diseases (ICD-10) is different from its predecessor, ICD-9, in several ways. In addition to a revised content that reflects the most important recent advances in research and clinical practice, it is presented in different versions for different types of professional users. The differences are, however, in degrees of detail and the versions are compatible with each other, since they are all derived from the same basic document (*Clinical Descriptions and Diagnostic Guidelines*, WHO, 1992)¹.

THE USES OF ICD-10

ICD-10 has necessarily retained its historical purpose of facilitating the recording of national and international statistics of morbidity and mortality, but now has the added values of also being designed as a uniquely international aide to clinical work, teaching and research. It achieves these objectives by means of an updated list of diagnostic rubrics, a set of glossary-type definitions of disorders and additional explicit diagnostic criteria. The latter have been developed in two versions: (a) clinical diagnostic guidelines for routine use, allowing sufficient flexibility and discretion in the application of "clinical judgement" in the hospital ward or the outpatient service; and (b) diagnostic criteria for research (ICD-10-DCR), providing stringent decision-making rules to increase the specificity of diagnostic classification and thus ensure a high level of sample homogeneity for the purposes of clinical, biological and other research².

As a result of a great deal of collaboration between the advisers to the World Health Organization and the several Task Forces that assembled DSM-IV on behalf of the American Psychiatric Association during the last few years of the preparation of both the classifications, there are very few important differences between them. Since the same body of internationally published research experience and literature was available to both sets of experts during the processes of development, those differences that remain are mainly differences of opinion rather than of fact. Some differences reflect the need for ICD-10 to accommodate a much broader base of international experience and opinions than a national classification. In the development of ICD-10, experts from many different cultures and languages were involved from the earliest stages.

As in ICD-9, Chapter V deals with "Mental and Behavioural Disorders" and is intended for the recording of the clinical syndromes as presented and experienced by the patient. If a specific underlying cause of the disorder is known (or highly

probable), additional codes should also be used from other chapters of ICD-10, such as: Chapter I, Infectious and Parasitic Diseases; Chapter II, Neoplasms; or Chapter VI, Diseases of the Nervous System.

DEMENTIA IN ICD-10

In ICD-10, the dementias are embedded in the section on organic and symptomatic mental disorders (codes F00–F09), which contains the following major rubrics:

- Dementia in Alzheimer's disease.
- Vascular dementia.
- Dementia in diseases classified elsewhere.
- Unspecified dementia.
- Organic amnesic syndrome, other than induced by alcohol and drugs.
- Delirium, other than induced by alcohol and drugs.
- Other mental disorders due to brain damage and dysfunction and to physical disease.
- Personality and behavioural disorders due to brain disease, damage and dysfunction
- Unspecified organic or symptomatic mental disorder.

In contrast to ICD-9, the distinction between psychotic and non-psychotic illnesses is of no taxonomic consequence in ICD-10, where disorders of different psychopathological expression are grouped together on the basis of established or presumed common aetiology. In the particular instance of section F0, in which the dementing disorders are included, the underlying classificatory characteristic of "organic" is defined in the sense that "the syndrome so classified can be attributed to an independently diagnosable cerebral or systemic disease or disorder". The subsidiary term "symptomatic" is not used in the titles of individual disorders but it is included in the overall title of the block F00–F09. This is because it is widely used in many countries to indicate those organic mental disorders in which cerebral involvement is secondary to a systemic extra-cerebral disease or disorder. In other words, "symptomatic" in this context is a subdivision of the wider term "organic".

Another feature of ICD-10, as compared to earlier classifications, is the omission of any reference to age as a defining criterion of the disorders accompanied by a cognitive deficit. The terms "senile" and "presenile" are practically absent in the classification, and there is no provision for identifying any mental disorder as necessarily a result of ageing. The classification does, however,

allow the recording of an unusually early or late onset of the disorder. In other words, the mental disorders occurring in the elderly are no longer considered to belong in a separate category of morbidity. This is very much in line with research conducted in the past two decades, which has demonstrated that the relatively high prevalence of mental morbidity in the elderly in Western cultures is related to a wide range of psychosocial factors (e.g. social isolation, cultural uprooting and institutionalization), as well as to physical co-morbidity, but that the aging process itself does not produce nosologically specific forms of disorders.

If section F0 of ICD-10 is used as a diagnostic decision tree, there is a choice of five entry points at the level of clinical syndrome: (i) dementia; (ii) amnesic syndrome; (iii) delirium; (iv) organic quasi-functional disorder (affective, delusional, hallucinatory or other); and (v) personality or behavioural disorder. Once a disorder is identified at this general syndrome level, the next step is defined by the diagnostic guidelines, which lead into more specific diagnostic categories. The diagnostic decision rules for dementia illustrate the point.

The syndrome of dementia is defined in ICD-10 by "evidence of a decline in both memory and thinking, which is of a degree sufficient to impair functioning in daily living", in a setting of clear consciousness. For a confident diagnosis to be established, such disturbances should have been present for at least 6 months. Deterioration of emotional control, social behaviour and motivation represent significant additional features but the overriding criterion is the presence of memory, learning and reasoning decline. The ICD-10-DCR (research criteria) add anchor points for a grading of the deficits into mild, moderate and severe, separately for memory and intellectual capacity. The overall grading of the severity of dementia is made on the basis of the function which is more severely impaired.

Once the presence of the syndrome of dementia is established, the diagnostic process branches off into the different clinical varieties of dementia typical of Alzheimer's disease, vascular dementia and dementia in diseases classified elsewhere (including Pick's disease, Creutzfeldt-Jakob disease, Huntington's disease, Parkinson's disease, HIV disease and a range of systemic and infectious diseases, such as hepatolenticular degeneration, lupus erythematosus, trypanosomiasis and general paresis). Dementia in Alzheimer's disease is subdivided into Type 1 (onset after the age of 65) and Type 2 (onset before the age of 65). Although the ICD-10-DCR criteria emphasize the ultimate criterion of the neuropathological examination and the supporting role of brain imaging, they nevertheless allow for a confident clinical diagnosis to be made if clear evidence of a memory and intellectual performance deterioration has been present for 6 months or more. The ICD-10 criteria for vascular dementia are broader than the corresponding DSM-IV criteria: they include not only multi-infarct (predominantly cortical) vascular dementia but also the subcortical dementias (Binswanger's encephalopathy being an example), as well as the mixed cortical and subcortical forms.

As regards the diagnosis of delirium, ICD-10 has abandoned the distinction between acute and subacute deliria; the condition is defined as "a unitary syndrome of variable duration and degree of severity ranging from mild to very grave", with an upper limit of 6 months' duration and a subdivision into delirium superimposed on dementia and delirium not superimposed on dementia.

The rubric "other mental disorders due to brain damage and dysfunction and to physical disease" includes disorders with "functional" characteristics (e.g. hallucinosis, catatonia, schizophrenia-like disorder, and mood disorders) that arise in the context of demonstrable organic illness, such as cerebral disease, systemic disorders and brain dysfunction associated with toxic disorders (other than due to alcohol or drugs). An important, not yet fully validated, addition to this rubric is the "mild cognitive disorder" attributable to physical co-morbidity (including HIV

disease), which is defined as transient but nevertheless involving memory and learning difficulties.

Finally, personality and behavioural disorders due to brain disease, damage and dysfunction include familiar conditions such as organic personality disorder (the frontal lobe syndrome, but also other lesions to circumscribed areas of the brain), postencephalitic syndrome, postconcussional syndrome and some new entities, e.g. right hemispheric organic affective disorder (altered ability to express and comprehend emotion without true depression).

In conclusion, two features of ICD-10 should be emphasized. First, as already noted, it does not identify the mental disorders in the elderly as a separate or special category of psychiatric morbidity. In addition to the F0 section listing the organic and symptomatic mental disorders, psychiatric disturbances arising in the elderly population can be classified, according to their presentation and course, in any of the other major sections of ICD-10 (except for F8 and F9, which deal with developmental disorders and behavioural and emotional disorders occurring in childhood and adolescence).

Second, although ICD-10 is not explicitly a multi-axial classification, there are two ways in which multi-axial coding can be achieved if required. The simplest way is to use extra codes from the other chapters of ICD-10 in addition to those in Chapter V; any physical disorders present can be recorded by codes from Chapters I–XIX, and codes from the final two chapters can be used to cover other noteworthy aspects of the clinical picture. These are: Chapter XX, External Causes of Morbidity and Mortality (the X and Y codes, covering drugs causing adverse effects in therapeutic use, and injuries and poisoning); and Chapter XXI, Factors Influencing Health Status and Contact with Health Services (the Z codes, which include a variety of social, family and life-style factors). Another and more comprehensive option is to use the special Multi-axial System now available, which was developed by means of an international collaborative study organized by WHO Geneva³. This provides three descriptive axes: Axis I, Clinical Diagnosis; Axis II, Disablements; and Axis III, Contextual Factors. These Axes are a convenient re-arrangement of the chapters of ICD-10 listed above, with the addition of a brief set of ratings covering physical disabilities.

PROSPECTS FOR THE FUTURE

It is likely to be many years before the next edition of the international classification is ready for use, but meanwhile there are plenty of issues worthy of debate. Whatever form the classification takes, it seems likely that the principle of recording the clinical picture by means of Chapter V and the underlying physical cause by means of other chapters will remain. Research is now providing many clues about the exact place and the histological nature of the lesions in the central nervous system that give rise to the clinical syndromes, but the clinical syndromes themselves will not change, and will always need to be recorded.

The new non-invasive techniques for brain imaging, such as MRI, SPECT and PET scans, are demonstrating a variety of structural abnormalities in the brains of some (but by no means all) patients with the familiar clinical syndromes that are also present in substantial proportions of normal subjects. These changes are "organic" in a general sense of being something physical, but not in the way the term is used in the ICD (that is, to indicate a concurrent and diagnosable physical disorder). It will probably soon be time to abandon "organic" as a term to be used in a classification, and to develop new terms and concepts that will make these more subtle differentiations clear.

It should be possible to omit the "nested" categories of organically caused syndromes of depression, anxiety and

schizophrenia in future versions of Chapter V. Their presence breaks the rule (otherwise followed) that a clinical syndrome should have only one place in the classification. They were included because of a clinical demand for the convenience of being able to specify both a syndrome and its cause by means of only one code. Their omission would cause no loss, while improving the properties of the ICD as a classification. Even desk-top recording systems now have large capacities and increasingly sophisticated software that make them capable of handling many entries for each clinical encounter or spell of patient care.

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Psychiatric Diagnosis and Old Age: New Perspectives for “DSM-IV-TR” and Beyond

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Aging and old age confront the psychiatric clinician and nosologist with special diagnostic problems. Our present syndromology has arisen largely from studying disorders of young and middle-aged adults. It is being challenged now by the general aging of the population in Western as well as many Eastern societies.

Aging reflects evolving biological, psychological, and social processes. Fundamental alterations of central nervous system functioning color the psychopathological and pathophysiological significance of specific symptoms, while the psychosocial meaning of discrete life events changes as one moves across the age spectrum. Old age brings an increase in confounding systemic medical conditions. Thus, we are confronted with an array of yet-to-be-answered questions. What is the relationship between the mentally disordered who have grown old, and the old who develop mental disorders? Does later age of onset connote a fundamentally different disease process, even when the presenting psychopathology is generally similar? How do we separate idiopathic (called “primary” in DSM-IV) psychiatric syndromes from psychopathological conditions caused by defined disease processes? What must the diagnostician and treating clinician do to distinguish those psychopathological symptoms that are amenable to treatment from confounding medical symptoms that reflect systemic illness?

The development of the fourth edition of the American Psychiatric Association’s Diagnostic and Statistical Manual (DSM-IV) provided an opportunity to begin considering such issues more formally. These types of questions were almost neglected in previous editions of the DSM, in part due to a lack of meaningful research data, and no doubt reflecting a lack of interest among many American psychiatrists in treating geriatric patients. However, the past 15–20 years have seen a surge in interest in the USA in psychogeriatrics, as evidenced by enhanced clinical training and sharpened clinical identity. The development by the American Board of Psychiatry and Neurology of an “added qualification” in geriatric psychiatry during the past decade crystallized this recognition.

There were relatively few changes in DSM-III or DSM-III-R that pertained to elders. Dementia and delirium were defined more precisely. The term “involutional” was dropped from the classification, due to a paucity of data supporting the qualitative distinctness of involutional melancholia. A criterion had been added in DSM-III to establish a maximum age of onset for schizophrenia at 45 years, but this was deleted in DSM-III-R as more evidence was made available to American writers that

schizophrenia-like disorders emerge among older patients and are not solely caused by primary cerebral diseases.

Literature reviews and discussions that were part of the preparation of DSM-IV revealed that there was a continuing lack of rigorous clinical descriptive research in many areas of geriatric psychiatry. There was a dearth of data regarding the natural history of both late-onset disorders and early-onset disorders that persist or recur throughout the life course into old age. It was also apparent that there were insufficient epidemiologic findings to fully define the prevalence or incidence of many later life disorders. In the USA, this may have reflected, in part, an overly rigorous application of DSM-III criteria during the epidemiological catchment area (ECA) studies. Although these were valuable for describing the psychopathology of some elderly patients, the stereotypic use of diagnostic descriptors developed for younger patients had the unintended effect of excluding possible subjects from each of the rubrics. This process was compounded further by employing lay interviewers asking highly structured questions. Many of the critiques regarding traditional approaches to cross-cultural psychiatry (e.g. the problem of “category fallacy”¹) are equally germane to studies that utilize criteria developed for one age cohort when characterizing the psychopathology evidenced by another. As a result, many recommendations for DSM-IV regarding later life psychiatric disorders were qualitative or impressionistic, more useful for specifying directions for new research but insufficient for substantially revising many of the diagnoses to be included in that new edition.

DSM-IV AND BEYOND

The major innovation of DSM-III was the development of a multiaxial system that provided clinicians with a wider array of descriptive categories for more completely defining their patients’ problems. However, there was little evidence in either DSM-III or DSM-III-R that indicated a thorough understanding of the complex multidimensional diagnostic problems posed by aging patients. In contrast, DSM-IV began the process of addressing this deficiency by including for many diagnoses a discussion labeled, “Specific Culture, Age, and Gender Features” or “Specific Age and Gender Features”. This approach allows some limited consideration of aging-related issues, but again, the sparse commentary in many of these sections also underscores how little is known about many related questions.

The following discussion reviews each of the DSM-IV axes, considering aging-specific topics. They point to areas of controversy that will need further study before inclusion in a future DSM-V. As well, they will highlight planned alterations for the forthcoming text revision of DSM-IV ("DSM-IV-TR") with regard to the diagnosis and classification of neurodegenerative processes, such as Alzheimer's disease.

Axis I

DSM-III and DSM-III-R continued the tradition of distinguishing between "organic" and "functional" disorders. This conceptual typology had posited a difficult-to-maintain dichotomy, one that was poorly justified in light of modern research². As well, "organic" diagnoses too often had been applied without any clear rationale. Instead, DSM-IV emphasizes a conceptual move away from concerns about the presence of structural pathology, or its absence, to a consideration of etiologic link.

It is more meaningful, both clinically and heuristically, to label a disorder as "primary" (i.e. idiopathic) when its cause is unknown, rather than using a term such as "functional" that implies a specific type of physiological or psychological mechanism. "Organic" arose in the nineteenth century after the detection of organ (i.e. brain) pathology. Research findings of the past 30 years have clouded any hope of using such a criterion for separating alleged psychological from alleged biological disturbances. DSM-IV expects clinicians to clarify whether a disorder is primary (i.e. idiopathic) or "secondary" or "symptomatic". (The text itself avoids using the latter designations, given controversy in the USA regarding their exact definitions.) A lengthy description is provided in DSM-IV to help clinicians reason whether there is sufficient evidence to attribute the etiology of a psychiatric syndrome to a definable disease process. Thus, one might diagnose "mood disorder due to Parkinson's disease" or "psychotic disorder due to Huntington's disease" where there is a decision that the psychopathology is a direct reflection of the fundamental pathophysiological process that causes Parkinson's disease or Huntington's disease.

While DSM-IV includes a chapter describing symptomatic psychiatric conditions, they are distributed throughout the text, grouped with the primary psychiatric disorders with which they share phenomenological features, forcing clinicians to consider differential diagnostic questions directly. Although the impact of this change extends beyond older patients, it has frequent application to elders, given their heightened rates of general medical conditions. Whether geriatric psychiatrists and other clinicians have, in fact, undertaken a more rigorous approach to considering etiological relationships because of the changes in DSM-IV remains to be defined.

Cognitive Disorders

Delirium, dementia and amnesic disorders were clustered together in DSM-IV as "cognitive disorders". DSM-III-R developed, perhaps inadvertently, very restrictive criteria for delirium that excluded a variety of patients whose clinicians were certain about the diagnosis, but unable to use the diagnostic label. Beyond traditional definitions employing attentional dysfunction and fluctuating consciousness, DSM-III-R required the presence of "disorganized thinking", as evidenced by formal thought disturbance. However, mildly delirious patients may not have rambling, irrelevant or incoherent speech. DSM-IV returned to a broader view.

While the successive editions of the DSM have been the primary classification for American psychiatry, ICD-9-CM (*International*

Classification of Diseases, 9th edn, Clinical Modification) has been adopted by international treaty as the medical diagnostic coding system used by the US Government for all data collection and reporting regarding morbidity and mortality. In order to receive reimbursement from Medicare, for example, every clinician must report diagnoses in terms of ICD-9-CM codes. These range from 00–0999, and are organized in sections based upon the type of disease process (e.g. infectious and parasitic diseases, 001–139; neoplasms, 140–239), anatomic location (e.g. diseases of the digestive system, 520–579; diseases of the nervous system and sense organs, 320–389) or functional area (e.g. mental disorders, 290–319).

The entire system of diagnostic coding now is being revamped during the development of ICD-10-CM. This is intended to deal with the problem of "double coding" (e.g. diagnosing meningitis as both an infection and a disease of the nervous system) and other shortcomings, most notably an insufficient number of potentially available codes for future classification needs. Beginning with the completion of ICD-10 by the World Health Organization during the early 1990s, the most important change has been the adoption of an alpha-numeric coding system, in which letters are used to indicate the section of the ICD classification from which the code is drawn. Thus, infectious disease codes start with the letter A, neoplasms start with the letter B, etc. Codes formerly contained in the Mental Disorders section (290–309) will start with the letter F, while codes formerly contained in the Diseases of the Nervous System (320–389) will be included under the letter G. The "clinical modification" of ICD-10 currently being developed by the US National Center for Health Statistics, using this alpha-numeric system, will be adopted officially within the next several years. The most significant change for geriatric psychiatry will be the provision in ICD-10-CM of only one code for Alzheimer's disease, G30, instead of the double coding that was part of DSM-IV and ICD-9-CM. Although sufficiently similar to ICD-10 to allow cross-national comparison of health data, ICD-10-CM will have several significant differences in terminology and level of specificity.

Subtypes of dementia of the Alzheimer type (DAT) and vascular dementia (i.e. with delirium, with depressed mood, and with delusions) were first introduced into the DSM-III at a time when these categories were referred to as "primary degenerative dementia" and "multi-infarct dementia". These subtypes were later carried forward into DSM-III-R and DSM-IV. The expressed rationale for adding these subtypes had been parsimony of diagnosis, based on the assumption that these three conditions were fundamental manifestations of dementia. It did not make sense to require a clinician to use an additional diagnosis to describe what were considered to be aspects of the primary clinical condition. But neither DSM-III nor DSM-III-R provided explicit instructions about when to use specific subtypes, and a number of questions arose regarding their use. For example, did "with depressed mood" include full-blown major depressive episodes occurring during the course of Alzheimer's disease, or just milder forms of depression? Should "with delirium" include all forms of superimposed delirium, or just delirium thought to be directly due to Alzheimer's disease or vascular pathology? How should one diagnose delirium secondary to suspected drug toxicity in a demented patient? What should be one's approach to describing other commonly encountered symptoms or signs, e.g. delusions, hallucinations, agitation, or anxiety? When delirium, delusions and depressed mood occur simultaneously, which diagnostic code does one choose?

DSM-IV continued to use this subtyping with three changes. A new subtype, "with behavioral disturbances", was added in order to allow the user to indicate "clinically significant behavioral changes (e.g. wandering)", although this was not codable in any numeric fashion. The "with delusions" subtype

required that the delusions be the “predominant feature”. Lastly, DSM-IV clarified that depressive presentations meeting criteria for a major depressive episode should be included under the “with depressed mood” subtype, in addition to milder forms.

Even with these changes, the basic reasoning behind the DSM-III, DSM-III-R and DSM-IV subtypes was flawed. Beyond including only a small portion of the array of behavioral, psychological or emotional signs, symptoms or syndromes that can be caused by Alzheimer’s disease, the structured hierarchy of the classification implied that these manifestations were a *secondary* or *subordinate feature* of the cognitive symptoms and signs of dementia. Ample data now demonstrate that, when present, they are most often manifestations of fundamental brain pathophysiology, as much as memory or other cognitive dysfunctions. In sum, for a classification that was intended to guide therapeutics as well as communicate information regarding signs, symptoms and prognosis, the system did not work.

A change in coding adopted by ICD-9-CM rendered these subtypes obsolete. In anticipation of planned changes for ICD-10-CM, the October 1997 ICD-9-CM coding recommendations indicated that the preferred diagnostic code for Dementia of the Alzheimer’s Type should be 294.1 (for “Dementia in Diseases Classified Elsewhere”) instead of 290.xx. Thus, the three subtypes that had been coded with a fifth digit could no longer be captured. These changes are being used to revise the approach in DSM-IV as well.

When a patient has emotional, mood or psychological symptoms or signs that are in need of therapeutic interventions arising from Alzheimer’s disease, the clinician will be asked to code these conditions on Axis I as one of the “mental disorders due to a general medical condition”. As with other etiologically- or pathologically-defined medical conditions, Alzheimer’s disease will be recorded on Axis III. Like other secondary or symptomatic psychiatric conditions, these problems must substantially interfere with a patient’s functional integrity to warrant a formal diagnosis.

Beyond dementia itself, the secondary conditions to be coded on Axis I (with an associated Axis III diagnosis of Alzheimer’s disease) include psychotic disorder, mood disorder, anxiety disorder, personality change (types include labile, disinhibited, aggressive, apathetic, paranoid, other, combined, and unspecified) and sleep disorder. As there are no data indicating that AD *causes* delirium, despite the frequent occurrence of delirium among patients with dementia of the Alzheimer’s type, the clinician will be pressed to define the specific etiology of the delirious condition when possible. Sexual disorders due to AD also should not be diagnosed, again reflecting an absence of clinical or research findings tying Alzheimer’s disease to sexual dysfunction. Those patients who are uncontrolled in their sexual behaviors can be captured under the “with behavioral disturbance” subtype. Thus, for a man with AD having dementia and delusions associated with combative behavior, one would use the diagnoses 294.1 and 293.81, the latter for Psychotic Disorder due to Alzheimer’s Disease, with Delusions. For a woman with delirium superimposed on DAT, the diagnosis will reflect the presumed etiology of the delirium (e.g. delirium due to anticholinergic use is diagnosed 292.81, Anticholinergic-induced Delirium).

Starting in October 2000, a new fifth digit will allow one to indicate whether the dementia is “with behavioral disturbances” (294.11) or “without behavioral disturbances” (294.10). The inclusion of a codable behavioral descriptor and the conventions regarding the diagnosis of DAT and other symptoms due to Alzheimer’s disease will be included in DSM-IV-TR, with an anticipated publication in May 2000. (The DSM-IV text revision project is an empirically-based updating of the text only, without any changes being made to the criteria sets.)

The term “*behavioral and psychological symptoms of dementia*” (BPSD) has not been adopted during the DSM-IV revision process. There are two explanations. The first relates to the label itself. It implies that the signs and symptoms in question are a direct outgrowth of dementia, which is not a fundamental disease process but itself a secondary clinical syndrome, rather than suggesting that the behavioral and psychological symptoms are due to AD-caused brain degeneration. The second reservation reflects a concern that behavioral, psychological and emotional symptoms and signs are not a unitary phenomenon. It is most parsimonious at this time to use the extant, discretely defined “mental disorders due to . . .” and avoid creating yet another unitary diagnostic label that has no established reliability or validity⁴.

A major topic for discussion during the development of DSM-IV was a recommendation that “age-associated memory impairment” (AAMI) be included as a diagnostic entity, in keeping with criteria proposed to standardize the definition of aging-associated changes in intellect⁵. Aging-related changes in intellectual ability are robust and demonstrable psychometrically when comparing healthy older subjects with younger control groups, encompass a variety of tasks beyond those related to memory, and are at times troubling for particular individuals. However, doubts were expressed regarding the use of a disease diagnosis for a normative phenomenon. There were no objectively defined standards for establishing a cut-off or threshold to separate it from the earliest manifestations of specific diseases (e.g. the early manifestations of Alzheimer’s disease). Indeed, review of available data suggested that mild intellectual declines *relative to age-matched peers* often presaged the development of progressive disease. In contrast, there was clear evidence that AAMI was *not* associated with significant functional or social impairment. Ultimately it was concluded that there were insufficient data to establish a formal psychiatric diagnosis of AAMI⁶. DSM-IV finally included “age-related cognitive decline” as a Z code designation, one of those conditions “not attributable to mental disorders that are a focus of attention or treatment”.

Other Axis I Issues

Other Axis I disorders provided focal points for heated debate, but concrete recommendations for classificatory changes were difficult to establish in the absence of well-developed research findings. The problem of mood disorders was illustrative. DSM-III-R criteria amply described many of the features of severe mood disorders seen in elderly patients coming to outpatient clinics or hospital inpatient services. However, they were far less satisfactory for describing the features of affectively impaired patients encountered in the offices of primary care physicians, evaluated in nursing home settings or ascertained through community surveys.

Many suffer “subsyndromal” presentations, either dysthymic states that generally conform phenomenologically to dysthymic disorder but are not sustained for the requisite 2 years, or manifestations that include mood, ideational or somatic features of an affective disorder but without the array of symptoms needed to qualify for a strictly defined diagnosis. A diagnostic system that excludes the majority of potential patients fails to fully serve its descriptive function⁷. But suggestions for change remain incomplete and are not yet supported by a sufficiently large body of careful clinical research. (So-called “minor depression” was added for research purposes to the Appendix of DSM-IV, but it remains uncertain how to apply this construct to the conditions encountered among elders.)

The deletion from DSM-III-R of age 45 years as a cut-off for the onset of schizophrenia removed a major impediment for

diagnosing elderly psychotic disorders. However, many clinicians have found unreliable the attempt to distinguish between schizophrenia and delusional (paranoid) disorder, based upon the presence of “bizarre delusions”. There is no standard by which one can decide when a delusion becomes “bizarre”. Other grounds for distinguishing delusional disorder from schizophrenia are now being considered. Additionally, current criteria for schizophrenia require a significant functional decline during the course of the disturbance, but investigators have noted that many patients with late-onset psychosis maintain stability in their personal and social functioning.

It is rare to encounter elderly patients with pure generalized anxiety disorders. Mixed anxiety–depression is common, but it does not fit easily among mood disorders or among anxiety disorders. Although some continue to advocate consideration of a label such as “mixed dysphoric state”, there has been no consensus. Growth in the use of serotonin reuptake inhibitors, and compounds such as extended release venlafaxine, has dampened the debate in practical terms, but their apparent effectiveness for mixed anxiety–depressive states sheds little light that clarifies the nosological confusion.

Regarding obsessive–compulsive disorder, post-traumatic stress disorder and phobias, there are few data available about their occurrence among elders to make any salient recommendations regarding diagnostic classification or modification based on aging-related changes in prevalence, presentation, course, co-morbidity or treatment responsiveness. In a similar vein, there are only a small number of replicated findings regarding the form or frequency of adjustment disorders among older patient groups. However, it has been amply clear that the currently employed 6-month time limit for adjustment disorders is inadequate when one faces a persisting stressful situation, such as a chronic physical disability or the need to care for a spouse with Alzheimer’s disease.

While there were some minor text changes in DSM-IV regarding substance use disorders, there has continued to be relatively little attention to specific issues related to older persons. Recent studies⁸ have underscored the variety of misconceptions that clinicians hold regarding the frequency and impact of alcohol-related clinical conditions. Many substance abuse problems among elders arise from misuse of prescribed medications; older patients often avoid the adverse social consequences of drug seeking and the medical complications of illicit intravenous administration. Although the abuse/misuse may be physically hazardous, especially as it relates to changes in metabolism or potentials for drug interactions, it is distinct from the jeopardy experienced by younger abusers. Thus, the diagnostic classification must attempt to take into consideration age- and culture-related variations, with an eye to dealing with unsupervised use as well as frank abuse.

Axis II

Consideration of personality factors and related clinical disorders has been hampered by minimal data. Although one may conjecture about the changing presentation of Axis II psychopathology across the age span, there have been remarkably few relevant, systematic studies⁹. Clinicians often recognize residual dysfunctional personality features in older, previously diagnosed patients who later fail to conform to stereotypic descriptions. There are no categories available for denoting such conditions. Similarly, there is no diagnosis of “emergent” personality disorder for describing those patients who, having suffered marginally impairing personality traits throughout their lives, evolve a frank disorder in old age. For example, such a classification would capture those who, having been supported or buffered by others,

become dysfunctional following the death of a spouse or parent. This view stresses the setting-dependent nature of disordered personality. Overall, pre-DSM-IV discussions swerved away from issues related to personality and aging, leaving them to be resolved in future editions. It is clear, however, that the domain of personality dysfunction—even if not captured by current diagnostic categories—is a major component of the geriatric landscape⁹, especially when considering problems such as depression in primary care settings^{10,11} or suicide in elders¹².

Axis III

As noted previously, DSM-IV includes guidelines for assisting a clinician in discriminating between a primary Axis I disorder and a symptomatic disorder due to a condition diagnosed on Axis III. When there is an etiologic tie, the general medical disorder should be noted as part of the Axis I diagnosis (e.g. “mood disorder due to Alzheimer’s disease”), in addition to its notation on Axis III. Despite the presence of guidelines, no rule can be used as an infallible aid to determine whether a condition is truly a manifestation of a fundamental medical disorder or is an unrelated idiopathic (“primary”) presentation that occurs by coincidence in the presence of a medical condition that is not tied etiologically to the Axis I psychiatric disorder. Clinical judgment must prevail. Similarly, there are no means available for indicating the current clinical significance of co-morbid conditions, whether they contribute to the patient’s overall disability, are confounds of possible treatments, or are merely incidental, coexisting disturbances that have no therapeutic impact or functional implications when considering the primary psychiatric syndromes.

Axis IV

As constituted in DSM-III-R, Axis IV was often unsuitable for use with elders. Its exemplars failed to account for many of the common problems or stresses of later life. Additionally, there was no method provided in the manual for defining the contribution of positive psychosocial factors that mitigated or diminished the contribution of stressors to the development of psychopathology. DSM-IV took another approach, defining what might be called “problem areas”. This approach avoided the shortcomings of the previous scale, and included a variety of domains relevant to elders, but it too provided no room for an assessment of compensatory factors, in addition to its problem-definition focus. A psychosocially orientated scale (or set of scales) is necessary, one that reflects a broad conception that includes social supports, occupational and environmental resources, perceived quality of life, as well as stressors, both acute and chronic.

Axis V

Functional ratings may have substantial predictive validity when used with elders, but also have proved unreliable when not used carefully. Current use of Axis V requires an estimation of functional capabilities based upon psychopathology alone. Many commentators question whether one can reliably separate functional deficits due to psychiatric symptoms from those arising from co-morbid physical disorders, especially in elderly patients with multiple diseases. Presently available global functional health measures are psychometrically robust and are excellent predictors of subsequent morbidity and mortality in older patient samples. Two separate rating scales, one devoted to overall global functional capability/disability and a second devoted to severity

of psychopathology, would likely show greater utility and reliability than the current Axis V measure. While this approach was discussed for DSM-IV, it was not adopted as it was viewed as untested. Future consideration of how best to rate a patient's functional abilities will be an important issue for DSM-V.

In a similar vein, development of a cognitive rating scale within the overall diagnostic classification would have substantial clinical and heuristic value. Here, too, one confronts substantial controversy. Does the clinician attempt to rate absolute capability or to assess change (i.e. decline)? While a measure reflecting intellectual integrity or decline would assist when considering a wide variety of disorders (including dementia, mood disturbances and psychoses), how should one account for variations in literacy, education, cultural background or country of origin? At present, no single scale has proved suitable for all circumstances. Before inclusion in any formal diagnostic system, such a measure would require substantial testing to ensure both its validity and its reliability.

CONCLUSION

Geriatric psychiatry is only now achieving in the USA the attention and status that it has held for many years in other countries. Historically, clinicians and researchers who have worked in this area have had little impact on the nosologies recorded in the diagnostic manuals of the American Psychiatric Association. This situation is changing dramatically. However, specific suggestions for revision bear only as much weight as the strength of the research base upon which they are built. The rapid growth in North America of research dealing with psychopathology of the elderly is a heartening development, one portending future changes in DSM-V that we may not have yet anticipated.

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History and Mental Status Examination

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A successful psychiatric evaluation in the elderly, including a comprehensive mental status examination, involves integrating components of both medical and psychiatric clinical models. The classic medical model pursues symptoms and signs in a problem-orientated fashion, seeking to match them with a single unifying diagnosis and then designing an appropriate treatment strategy leading to a cure. This approach is less useful when dealing with geriatric patients. With these patients, aspects of the psychiatric clinical model are emphasized, i.e. ongoing treatment rather than cure, and chronic rather than acute care given by an interdisciplinary team. This strategy leads toward maximizing and maintaining individual function and behavior. Geriatric care is involved less with disease and more with disability. This is the case whether the patient has a psychiatric or general medical problem; and treatment goals must address the situation of chronic care needs, in contrast simply to the “fixing” of an isolated problem. Complicated presentations of disease in this age group create problems most successfully approached from an interdisciplinary perspective. Many syndromes manifest themselves with symptoms common to psychiatry, internal medicine and neurology, illustrating the overlap of these disciplines in geriatric care. The unique and challenging aspects of interviewing the elderly requires developing an individualized, functional clinical examination. With this goal in mind, the medical history will be examined and utilized to illustrate how the different medical specialties interact to integrate the art with the science of medicine. A successful blending will result in obtaining the most clinically useful examination possible.

PURPOSE

Obtaining a valid and reliable medical history continues to yield a correct diagnosis in over 80% of clinical situations¹. This remains true in elderly patients, but accomplishing the task in this age group is often difficult and time-consuming. Perfecting the skills required to interview successfully an old, sick individual is invaluable in understanding an unusual disease process or avoiding costly mistakes in evaluation and treatment². The working, hands-on abilities necessary to effect a successful history fall within two separate realms: fact-finding (science) and empathy (art)³. In the first realm, traditional medical training is replete with teaching and experiences. On the other hand, tutelage yielding artful practitioners is much less universal; indeed, it is becoming a rare and treasured experience in a medical residency.

With a view in mind toward expanding the “art” realm, a new look at interviewing elderly people is indicated. The interaction between patient and interviewer becomes much more than the

process of collecting facts and related information to make a diagnosis. Gaining medical and social facts pertaining to the individual’s problems is a primary but not exclusive goal. Older adults often come to the physician with a complex agenda that may not be obvious by dwelling only on the chief complaint. Conducting the interview by this conventional fact-finding method may not be productive and actually may put one at risk for generating too narrow a differential diagnosis.

In addition to laying the foundation for a satisfying therapeutic relationship, the initial gathering of information should guide the first steps in evaluation. Logical, stepwise investigation of symptoms and signs remains the practical approach to any medical work-up. If this investigation yields nothing, the urge to perform more elaborate tests should be resisted and one should return to the history. Greater dependence on reviewing the history again with the patient is more likely to yield an appropriate plan. The “law of parsimony” is not as useful a guiding principle in the care of geriatric patients as it is in general medicine⁴. A single, unifying diagnosis may not explain satisfactorily a symptom complex responsible for a precipitous functional decline in an elderly individual, and continuing investigation frequently uncovers multiple causes.

An understood but often unstated purpose of the medical interview with old people is getting to know the patient. This requires interviewer empathy (not sympathy or emotional involvement), which may be of the highest importance in the patient’s view. Using the initial interview to explore personality and life experiences helps build rapport that is essential to subsequent interactions. Whether a patient’s problem is psychiatric or medical, the initial interaction and relationship that grows from it are vital to achieving the goal of proper diagnosis and treatment.

ELEMENTS

Although the components of the medical history in geriatrics do not differ from those in the practice of any other specialty, the order in which they are pursued and their relative importance may be unique. The classic approach of taking a symptom-directed chief complaint, history of present illness, past medical history, review of systems, social history and family history will produce a desired database from which to work. However, mechanically and swiftly following an unbending order or sequence of impersonal questions may provoke successively less and less cooperation if the patient perceives that you are not really “listening”. A more effective approach may require the physician to give up some control over the process of the interview, allowing the patient to

conduct it on his/her own terms. Valuable information that would otherwise be missed may be hidden in a conversational tangent the patient insists on following.

Review of previous medical records is necessary but they may not be available while you are seeing the patient. Especially in the elderly, a variety of information sources must be accessed in addition to the patient, including reliable family members and others in the social network. Indeed, if these other sources are lacking this should be noted and may itself have significance in the geriatric setting. Even in cognitively intact persons, corroboration is a good idea, as the "facts" may be related differently depending upon one's perspective. Non-medical problems, such as financial stress or social isolation, invariably impact on the elderly patient in unpredictable ways and cannot be ignored while investigating a medical complaint. After thorough investigation of each part of the history, the stories, details and facts are compiled, reviewed and evaluated, resulting in an in-depth, composite picture of the person in relation to the clinical issue at hand.

EXECUTION

For the older patient the first encounter with a physician is very important. Any subsequent interaction will be strongly influenced by the patient's initial impression, e.g. empathetic or distracted, rushed or friendly, interested or detached, confident or tentative. All things will flow from the stage set in the first few minutes of the interview. Therefore, serious thought should go into the initial 5 or 10 minutes of conversation. The physician must approach the situation with confidence, courtesy and the belief that some treatment plan can be developed, no matter how hopeless the situation may initially seem. Indeed, an optimistic stance is often useful when dealing with an elderly patient. The only thing on the physician's mind during the interview should be the patient and the problem at hand.

A skilled interviewer simultaneously questions, interacts (exchange of information), observes and, most importantly, listens. Proficient listening is an active exercise, requiring great concentration. Well placed periods of silence can be fruitful regarding observation or obtaining additional volunteered information, if the atmosphere is supportive and relaxed. It is not necessary to keep a dialogue going non-stop throughout the interview. Silence should be considered as a positive strategy. Sometimes, silent observation is extremely useful, especially with patients who seem to be actively hallucinating or exhibiting abnormal motor movements.

The examiner must function at many levels of interaction simultaneously, which can be physically draining. Making adjustments and modifications for the sensory and sometimes cognitive impairment extant in some patients adds to the interviewer's stress. These modifications include: sitting close; speaking slowly and loudly with frequent repetition; provision of ample time to allow for responses, which often have a prolonged latency; and even, when possible and necessary, use of sound amplifying devices⁵.

More active engagement and even providing verbal direction may be a useful and effective tactic during the interview. The personal style of the interviewer is an individual but important trait, and the patient must be convinced that it is genuine. This occurs if the interviewer is truly interested and able to suspend any distracting thought or action. Physical expressions of concern, including touching (e.g. hand holding) are often appropriate with particular geriatric patients and helpful in reinforcing positive rapport.

The best initial tactic is to focus on the chief complaint, but not be limited by it. Know the planned questions and format of the entire interview well, allowing for rearrangement of sections

during a particular interview to suit individual situations. This will allow a fluid order without interruption so that an interesting tangent may be followed easily. Open-ended questions may not be productive with the elderly, and if no progress is being made, asking focused, specific questions or using different, sometimes simpler, wording may succeed.

In spite of thoughtfully executed and patiently stated questions, contradictory answers may still be the end result. At that point resist a frustrated summary and probe further, pushing for elaboration. The contradiction will either be resolved directly with new information, or may itself provide new insight when it cannot be explained. Patience is required of both the examiner and patient, and it may be necessary to execute the complete interview in several parts, deferring some material to subsequent sessions.

Astute observation of the patient throughout the entire interaction may yield more information than a multitude of questions. The interviewer, acting as a receiver, uses all senses—eyes, ears and especially immediate feelings—to accept and interpret transmitted information. Much is learned by gazing in the face of an elderly person, e.g. presence or absence of eye contact, facial musculature, position of the head and general affect. Subtle body language, such as hand motion or lack of it, can suggest affective disorder, neurologic or medication problems, or perhaps substantiate complaints of arthritis.

Commenting on objects or memorabilia observed in the surrounding environment serves the dual purpose of providing the opportunity for light conversation and again for building rapport. Reminiscence of earlier times, when the patient was in greater control, can also be a valuable method of gathering data, in addition to putting the patient at ease and strengthening the therapeutic relationship. For the unusually quiet elderly person, try specific "ice-breakers", such as:

"Tell me the story that is your life . . ."

"What was it like to be alive when (some significant event) . . .?"

"When you were my age, how long did you think you would live?"

This particular tactic may even provide the interviewer with insight into cohort differences among different-aged elderly and their impact on disease presentation in individual elderly patients.

Certain problem situations present unique challenges when interviewing the elderly. Unavoidable physical limitations, such as fatigue, hearing or visual impairment and chronic pain, may limit the time of the interview itself or influence the amount of information gained. Cognitive problems, such as memory loss or inability to pay attention, and psychiatric illnesses (temporary or permanent), such as dementia, delirium, generalized anxiety, major affective disorder or even psychosis, may cause difficulties and such diagnoses may only become apparent well into the interview, making all information obtained up to that point suspect. Also, patient biases of many types, such as racial, language and cultural, need to be acknowledged to avoid misinformation and miscommunication. Old patients may view young physicians with distrust, just as age bias may occur in the opposite direction.

Closing the interview successfully is as important in overall impact as is the initial impression⁶. Take care to ensure that all issues on the patient's agenda have been addressed. In any event, a useful strategy is always to end the interview by asking the patient if there are any questions he/she would like to ask the interviewer. Key questions may often be asked by the patient just as the physician closes the encounter, uncovering hidden fears or revealing the real reason (not the stated one) for the visit. Adequate time for patient questions, empathetic listening and thoughtful responses often yield insights missed or left unexplained earlier. A brief statement to the patient at the end of the

interview about “where to go from here” can instill confidence that something will be done soon. Later, after time for thoughtful consideration, a detailed explanation of the assessment and treatment plan can be discussed with all interested parties.

The primary goals of psychiatric and general medical treatment merge in the elderly patient⁵. They include the amelioration of symptoms, restoration or maintenance of optimum function, and enhancement of the subjective quality of life. Since curing the elderly of disease is usually not possible, focusing on these goals in a caring manner will enhance the physician–patient relationship and increase the chance of success in treatment.

Once the facts are gathered, one then needs to spend considerable time systematically reviewing and organizing the data into meaningful information, i.e. the formulation. This allows for the development of an integrated diagnosis and treatment plan. The plan is a framework and remains a guide, but can be altered depending on subsequent events and the patient’s response to therapy. A well thought out plan also helps avoid costly and often fruitless searching and testing that may lead to over-diagnosis and risky therapeutic choices. Both professional and personal rewards result when extra time and effort are invested in thoroughly examining an elderly person in this manner.

MENTAL STATUS EXAMINATION

The formal mental status examination (MSE) is obtained once a thorough history is obtained. It is a measure of both general psychiatric status and specific cognitive function or dysfunction. Equal time, patience and concentration are required for this portion of the interview, and, especially in the mentally ill elderly, often is a more difficult task for the interviewer. This is because the MSE touches on areas and experiences that may be uncomfortable, embarrassing or even frightening for the patient. Often, parts of the MSE will have already been satisfied by previous observation and interaction while taking the history, e.g.

level of consciousness, appearance and behavior, affect, and parts of the sensorium examination (attention, word-finding).

Table 24.1 displays the components of a formal MSE in the psychiatric interview. The patient’s level of consciousness should be described at the outset. Columns 1–4 of Table 24.1 are examples of common descriptors illustrating possible presentations of mental illness. The last four columns suggest areas to consider when investigating sensorium, judgement, insight and risk of causing harm.

At the conclusion of the MSE all acquired information can be recorded in a comprehensive psychiatric database, as outlined below. This serves many purposes, but most importantly it encourages systematic review, comparison with subsequent examinations, and decreases the possibility of overlooking details important to a correct diagnosis. Formulation of an individualized assessment and treatment plan, including functional status, completes the process and, once recorded, allows review and revision at any time. It should be reinforced that psychiatric evaluation of the geriatric patient, including history and MSE, although similar in some ways to examination of younger patients, differs in some key ways. The most important, unique difference for the examiner to understand and appreciate is the need to integrate into the MSE the assessment of concomitant chronic and acute medical conditions and also to augment the historical data obtained from the patient by interviewing other reliable individuals with pertinent information.

Database History

Identification

Informant(s)—determining reliability of the historian is important. It is often useful to supplement the interview by meeting and questioning family and/or friends

Chief complaint

History

Present illness

Past psychiatric

Table 24.1 Components of the mental status examination (MSE)

Appearance and behavior	Affect	Mechanics of thought (speech)	Content of thought	Sensorium (cognitive exam) ⁶	Judgment	Insight	Suicidal/homicidal
Grooming	Absent (blank)	(1) Quality:	Preoccupation	Orientation	Decision-making	Ability to understand	Ideation
Dress (gender-appropriate?)	Withdrawn	Rate	Obsession	Memory	capacity ⁷	important	Previous attempts
Mannerisms	Sad	Volume	Guilt	Registration	Health	components of a	Family history
Posture	Euphoric	Prosody	Somatic concerns	Recall	Financial	situation	Risk factors
Motor activity	Labile	Clarity	Paranoia	Long-term memory			Plan
Interviewer’s reaction to patient	Intense	Pressure	Erotica	Attention			
Patient’s reaction to interviewer	Constricted	(2) Quantity:	Delusions	Concentration			
	Flat	Spontaneous	Well-organized	Calculation			
	Appropriate to content of talk?	Mono- or polysyllabic	Bizarre	Abstraction			
			Hallucinations (sensory modality)	Comprehension			
				Consequences			
				Planning			
				Foresight			
				Reasoning			
				Problem-solving abilities			
				Language			
				Writing			
				Visual–spatial			

Disciplines other than psychiatry routinely test only orientation to satisfy this section of the MSE. It is important to perform a standard, valid and reliable cognitive examination to achieve an acceptably comprehensive MSE. The Mini-Mental State Examination (MMSE) is a good example of an acceptable test⁸. Also, besides testing cognition, other instruments may be useful in particular patients, especially to evaluate mood and functional capabilities⁹. The interviewer should become familiar and experienced with one or two scales in each area, e.g. Geriatric Depression Scale¹⁰ or Beck’s Depression Inventory¹¹; Activities of Daily Living¹² Instrumental Activities of Daily Living¹³; Global Deterioration Scale¹⁴; Cognitive Performance Test¹⁶ Global Assessment of Functioning¹⁷.

Family
 Family psychiatric
 Personal
 Social (including work)
 Medical
 Current medications
 Drug and alcohol
 Psychiatric examination (MSE)
 Level of consciousness
 Appearance and behavior
 Affect
 Mechanics (stream) of thought
 Content of thought
 Sensorium
 Insight
 Judgment
 Suicidal/homicidal ideation

A physical examination (including a brief neurologic examination), laboratory evaluation and neuropsychological testing complete the geriatric psychiatric database and are described elsewhere. It is particularly important to evaluate visual and auditory acuity in this age group.

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The Physician's Role

Lesley Young

The idea that mental disorders may be the result of bodily disease acting on the brain has been current for at least a millennium, since the time of Avicenna¹. Numerous attempts to relate specific psychological reactions to particular physical disorders have been unsuccessful, however, and it now seems clear that some general mechanism affecting the chemical environment of the brain is responsible². Many descriptive classifications of "organic psychosis" have had to be abandoned and not even the notion of "clouding of consciousness" has always been regarded as a cardinal feature of delirium³, although the most recent definition of delirium includes "disturbance of consciousness with reduced ability to focus, shift or sustain attention" as a diagnostic criterion⁴.

The clinician dealing with a mentally disturbed patient must consider any process capable of interfering with cerebral metabolism or neurotransmission^{5,8} as a possible causal factor. Moreover, for reasons still not well understood, the probability of a mental disorder arising from a "physical" cause is greatly increased in old age, so that the psychiatrist will rarely see a mentally ill old person without significant physical disease⁶ (and the geriatrician must likewise develop expertise in psychiatry).

The aim of the physician when confronted with a mentally ill patient should be to answer two questions:

- Could there be a physical cause for the mental disturbance?
- Is there any physical condition that is making things worse?

DELIRIUM

"Acute confusional states" are common in old people⁷, indeed it is claimed that mental confusion is a far more common herald of physical illness than pain, fever or tachycardias⁸. Delirium may also go unrecognized by attending clinicians⁹, or be misdiagnosed as dementia or depression¹⁰, with potentially disastrous consequences¹¹. Not only may this mean a denial of personal rights or even permanent incarceration in an institution, but also the underlying cause may go undetected and therefore untreated. Although detailed mental state examination and formal cognitive testing may give some clues, delirium and dementia can only reliably be distinguished by obtaining a full collateral history and by observation of the subsequent course of the illness. However, the serial use of cognitive screening tests, such as the Abbreviated Mental Test score¹² or Mini-Mental State Examination¹³, has been shown to be useful in helping to distinguish delirium from dementia. Disorders of attention and alertness are prominent features of delirium, and tests of reaction time and the "face/hand"¹⁴ tests have been claimed to distinguish those whose mental

disorder is of recent and rapid onset. Hallucinations and other perceptual disorders are also common and may be found in the absence of any other apparent mental illness, so such phenomena may be under-reported for obvious reasons. Visual hallucinations are said to be particularly common following bereavement¹⁵ and isolated perceptual disorders such as the Charles Bonnet syndrome¹⁶ may be associated with pathology of the sensory system involved, although such phenomena are also seen in cerebrovascular disease¹⁷. None of these features is unique to delirium, however, and the only reliable way to distinguish delirium from dementia, and sometimes from functional mental illness, is to obtain a detailed history from a third party: the most useful diagnostic tool is the telephone.

If in doubt it should be assumed that the mental disturbance is acute and therefore potentially reversible, so a physical cause should always be sought, a process that may require some detective work. Appropriately enough, the first step is to suspect that the victim has been poisoned. A full drug history, including both prescribed and over-the-counter medication, is necessary. Although drugs acting primarily on the central nervous system, and particularly those affecting the cholinergic neurotransmitter systems, are the most likely to cause problems, any medication should come under suspicion. Numerous drugs have been shown to have cholinergic effects, including many drugs that would not initially appear to exert such an action¹⁸. Polypharmacy is common in older people²⁷ and relatively minor cholinergic effects of each individual drug may exert a cumulative effect, thus, the delirium may not be due to one individual drug but rather the combined effect of several. Drugs affecting cardiac output or cerebral blood flow may precipitate memory changes, and diuretics also produce a range of adverse metabolic effects of their own. Long-term use of laxatives may have similar effects. "Minor tranquilizers", such as the benzodiazepines, once thought of as safe alternatives to the barbiturates, are now recognized to cause acute psychiatric disturbance¹⁹ as well as chronic mental impairment, unsteadiness and falls²⁰. Opiates, often prescribed together with mild analgesics in combinations such as co-proxamol, may cause acute or long-term problems.

Long-term use of addictive drugs is surprisingly common in the elderly population²¹ and is often the result of careless prescription of sleeping tablets or painkillers. Alcoholism is probably an under-recognized problem^{27,23} and in this context it should be remembered that delirium may just as easily result from sudden withdrawal of long-term drugs as from their use²⁴.

Examination of the patient often begins with inspection of the home, looking particularly for evidence of drug or alcohol abuse, poor diet, incontinence and general signs of short- or long-term neglect. The family and neighbours should be interviewed

wherever possible, mainly to establish the patient's normal mental and physical state and the timing of any change, making use of "landmarks" (such as Christmas) to focus the memory. The patient's current mental state should then be assessed in the light of this information.

Diagnosis and Assessment

When assessing an elderly person with a potential delirium, it is useful to bear in mind the common causes for precipitating delirium, which include infections, biochemical and metabolic derangements, organ failure and drugs. There is frequently more than one potential cause and thus assessment should be continued, even after one "cause" has been identified.

The approach and manner of the clinician towards the patient are of great importance. The aim is to provide a stable and, if possible, familiar sensory environment, so assessment should take place in a well-lit room with a minimum of distracting and misinterpretable stimuli. Physical examination starts with assessment of general hygiene, nutritional state, hydration and superficial signs of injury.

Body temperature must be measured with care: significant infections usually cause fever at any age²⁵, although it is often missed in elderly patients, where oral or axillary temperature may take longer to equilibrate with core temperature²⁶. Suspected hypothermia must be checked by measuring rectal temperature. The relationship between peripheral and core temperature depends mainly on the state of the circulation: cool peripheries are just as likely to indicate arterial shut-down, due to low cardiac output or hypovolaemia, as hypothermia. Other important signs of dehydration or blood loss are a low jugular venous pressure (venous pulsation not visible at the root of the neck with the patient lying at 20° or less to the horizontal) or postural hypotension (a fall of over 20 mmHg systolic and/or 10 mmHg diastolic on sitting or standing up). The latter may, of course, be due to the effect of drugs or autonomic failure, in which case the fall in blood pressure may not be accompanied by reflex tachycardia. High blood pressure has little diagnostic value and should normally be left alone in the acute situation.

Abnormalities of heart rate or rhythm can also seriously affect cardiac output and cerebral perfusion. Heart rate should be assessed by both feeling the pulse and listening over the precordium, where murmurs can also be detected. An electrocardiogram is an essential extension of clinical examination and, as well as documenting heart rhythm, may also show clinically undetectable signs of myocardial infarction. Signs of cardiac failure should be sought, including the presence of a gallop rhythm, fine inspiratory crepitations in the chest and peripheral oedema. The peripheral pulses should be felt. Bruits in the neck are useful, although non-specific, markers of arterial disease.

Abnormalities of respiratory rate or pattern must be noted, as well as the presence of central or peripheral cyanosis. In a delirious patient with cyanosis, warm peripheries and bounding pulses, the characteristic flapping tremor of CO₂ retention should be sought. Focal signs in the chest are useful but non-specific, and a chest X-ray is nearly always required. Any sputum must be examined at the bedside as well as sent for microscopy and culture. Arterial blood gasses are sometimes useful, but both persistent and transient hypoxaemia can be detected non-invasively with a pulse oximeter, although in the presence of signs of hypercapnia measurement of arterial blood gases is preferable.

Examination of the alimentary system begins with nutritional assessment and inspection of the mouth, looking for evidence of sepsis or neoplasm. The abdomen must be carefully examined, as

virtually any surgical emergency can present as a change in mental state without apparent abdominal symptoms. Abdominal signs may be far from obvious and many a strangulated hernia in an elderly patient has been missed by the unwary.

Urinary retention and faecal impaction are often detectable per abdomen but a rectal examination should also be done wherever possible. Whether constipation can cause delirium by itself is still a subject of vigorous debate, but most geriatricians and general surgeons are familiar with the elderly patient whose mental and physical state improves dramatically with rehydration and enemas. Examination of the perineum includes inspection of clothing for evidence of incontinence or precautions against it. Incontinence is a frequent finding in patients with delirium²¹; whether as a consequence of the underlying cause or as a direct result of the mental disturbance is not clear.

In the locomotor system, trauma and acute inflammation are important causes of delirium and swelling, warmth, tenderness and pain on movement of any joint should give rise to suspicion. The feet should be carefully examined for signs of ischaemia or sores, and the gait observed.

The most important part of neurological examination is the assessment of conscious level and higher mental function. In a delirious patient this is as much the province of the physician as the psychiatrist and it is important that both terminology and assessment should be standardized. The hard-pressed house physician confronted with an acutely disturbed patient in the middle of the night is unlikely to turn to the Diagnostic and Statistical Manual of Mental Disorders of the American Psychiatric Association for immediate guidance, but she might find a simple standardized assessment of attention, orientation and memory, such as the 10-question Abbreviated Mental Test Score²⁸ or the Mini-Mental State Examination¹³ useful. Such tests are often criticized because they are prone to misuse, a low score being taken as evidence of dementia, but their main value lies in repeated use, when dramatic changes often seen in acute medical patients clearly point²⁹ to delirium rather than dementia as the cause of poor performance. The routine use of such tests reminds the busy junior doctor to consider the possibility of delirium, and may prompt a search for the underlying cause in those who have a low score. Moreover, routine use of cognitive screening tests can avoid the clinician being fooled by an apparently cognitively intact older person who, in fact, has a significant cognitive deficit but a good "social front", can act as a baseline should the mental status change. It is important to be aware, however, that these tests are merely screening tests, and the scores are influenced by numerous factors other than the current mental disturbance. Including sensory impairments, educational level, social class and language. When applying standardized criteria to patients admitted to a geriatric assessment unit, 18% satisfied DSM-III-R criteria for delirium on admission²⁹ and the mean duration of the delirium by the same criteria was 7 days. Some deficits persisted longer, however, notably memory impairment, with a mean duration of 28 days. The use of standardized scales will not replace clinical judgement but will undoubtedly lead to greater clarity of thought and discussion, and is essential for progress in research.

Many clinicians place great emphasis on careful neurological examination, looking for focal signs which, if found are usually attributed to cerebrovascular disease. This assumption may often be correct, although the evidence is scant, but it is a mistake to blame everything on acute stroke, since the metabolic disturbance causing delirium may simply be highlighting areas of brain ischaemia. Thus, focal signs may be related to previous stroke damage, where neurological function has largely recovered although perfusion has remained impaired. If this is not appreciated, then the real culprit, which might be a treatable infection, may be missed.

The reliability of subtle focal signs, such as asymmetry in tendon reflexes, should also be regarded with scepticism, given the extent of inter-observer variation in the assessment of the patient with obvious hemiplegic stroke^{30,31}. The most reliable signs are those that can be easily reproduced and particularly those that are reflected in the patient's behaviour, such as dysphasia or unilateral neglect. This raises the more general point that the diagnostic assessment does not end with the formal physical examination but must include careful observation of the patient over the next few days. A host of signs that might otherwise have been missed, from episodes of fever, syncope, or fits to a craving for alcohol or opiates, may be revealed. Assessment of vision and hearing are particularly important, as the presence of sensory deficits may increase the risk of developing delirium, and a recent study showed that attention to and correction of sensory deficits may reduce the severity and duration of delirium³². Impairments of vision or hearing may also only become clear after a period of observation.

The choice of laboratory tests tends to be determined more by institutional tradition and by habits and attitudes of the clinician than by scientific evidence. The value of any tests depends largely on the prior probability of finding treatable disease. Thus, the urine should always be tested but a computed tomography (CT) scan comes low on the list of priorities. Whilst CT head scans may frequently show minor abnormalities, such as cerebral atrophy, a cause for delirium is rarely found unless there are other features in the history and examination to point to an intracerebral cause, such as new onset of focal neurological signs³³. Similarly, whilst electroencephalography (EEG) is frequently abnormal in delirium, showing diffuse slow-wave activity³⁴⁻³⁶, it is rarely useful as a diagnostic tool unless there is clinical suspicion of epilepsy. Indeed, the very non-specific nature of the EEG changes seen in delirium raises interesting (although under-researched) questions about the pathogenesis of the disorder. Other invasive tests, such as lumbar puncture (LP), have been used in research and numerous abnormalities of neurotransmitters have been identified in delirium, but as a diagnostic test in clinical circumstances, LP is again only useful in those with features of meningism^{37,38}.

The range of diagnostic possibilities is almost always wide and there may be few clinical clues. This has led to the erroneous and derogatory use of the expression "geriatric screening" to describe the laboratory investigation of elderly patients presenting with non-specific symptoms or functional problems. Screening tests always have a low yield, since the vast majority of subjects have nothing wrong with them, whereas tests done in order to track down the cause of delirium have a high probability of showing an abnormality, and serum calcium, urea and electrolytes, full blood count and glucose should virtually always be done.

DEMENTIA

Delirium and dementia commonly occur together; indeed, the pre-existence of dementia is a major risk factor for the development of delirium³⁹ and up to 70% of patients presenting with an episode of delirium will have some degree of chronic brain failure as well⁴⁰. There is a clear relationship between the degree of vulnerability of a patient and the size of the insult required to precipitate delirium³⁹. The principles of assessing a patient with "decompensated dementia" are similar to those outlined above, with some slight changes in emphasis. A psychological upset or disruption of the normal protective social environment may be enough to precipitate a crisis in a mentally frail person. Whether or not such an episode of decompensation actually constitutes delirium is debatable⁴¹ but it can only be a short step away, as physical complication such as exhaustion, dehydration or hypothermia may quickly supervene. Whether or not the initial

event was some kind of social or psychological trauma, it should not prevent a careful search for physical disease, for in such frail individuals, relatively minor remediable disorders can have dramatic functional consequences.

The distinction between decompensated dementia and delirium is also of practical importance because of the special problems involved in admitting the former group to hospital. A balance must be struck between the patient's need for a stable, familiar and reassuring environment and the availability of facilities for investigation and treatment. The former may be the overriding consideration in the case of decompensated dementia, while in the previously well person with delirium the latter is the main concern. In some (but not all) cases, day hospital facilities may provide a useful compromise.

Physical illness in a person with dementia may also present a decline in physical rather than mental function, with development of non-specific symptoms or signs such as unsteadiness, falls, immobility or incontinence. Again, the sudden appearance of such problems should not be assumed to be due to the progression of dementia or put down to "another stroke" but must be investigated, since timely intervention may prevent irreversible deterioration or even unnecessary institutionalization.

Recent research has highlighted the associations between physical illness and dementia, particularly vascular pathology and risk factors for such conditions. Dementia is thus significantly associated with the presence of atrial fibrillation⁴², ischaemic heart disease^{43,44}, cerebrovascular disease^{43,44}, hypertension^{44,45} and diabetes^{46,47}. Moreover, there is suggestive evidence that poor control in some of these conditions may result in accelerated cognitive decline⁴⁵. Therefore, whilst evidence from large-scale prospective studies is not yet available, it would seem sensible for the physician to actively treat such coexisting physical illness in the presence of dementia. Whilst high blood pressure appears to be a risk factor for the development of dementia, the prevalence of low blood pressure and orthostatic hypotension (OH) is also more common in dementia and may have an aetiological role^{48,49}, and the physician should be alert to the presence of OH and consider appropriate intervention.

Involving physicians in the assessment of patients with dementia is important to help rule out a treatable cause. Clearly no-one can afford to miss "reversible dementia" but unfortunately we have no idea of its true prevalence. Estimates of the frequency of reversibility in series of hospital patients has been reported to be as high as 40%⁵⁰, but older studies are hopelessly biased by concentrating on younger "pre-senile" patients from secondary or tertiary referral centres. In the few community-based studies that have been reported, the frequency of "reversible" cases was less than 10%⁵¹. More recent studies have estimated the prevalence of potentially reversible dementia in the region of 7.2%⁵² to 23%⁵³, although only at best 3% were actually reversed^{52,53}. An analogy may be drawn with the frequency of space-occupying lesions in patients presenting with "acute stroke", which was estimated at 15% in a study based at a neurological centre⁵⁴, compared to 1.5% in a series of patients admitted to an acute geriatric unit⁵⁵ and a similar figure in a community stroke survey⁵⁶. There is an urgent need for equivalent community-based studies of dementia in which all cases are thoroughly investigated. Until then, practice must be based on questionable and possibly ageist assumptions about the likelihood of finding treatable disease in particular groups of patients. Thus, few clinicians would disagree that younger patients should be fully investigated, as well as those of any age whose symptoms are of recent onset or rapidly progressing.

As emphasized above, a careful drug history is of overriding importance, since it is bad enough to miss a treatable disorder but unforgivable to be responsible for causing it or making it worse. As in the case of delirium, all drugs should come under suspicion,

including some superficially unlikely offenders such as digoxin, non-steroidal anti-inflammatories and cimetidine. The anticholinergic effects of phenothiazines, tricyclic antidepressants and anti-Parkinsonism drugs commonly cause problems. Longer-acting sulphonylureas may produce a state of chronic befuddlement in elderly patients, contrasting with the acute hypoglycaemic episodes and prominent adrenergic symptoms usually seen in younger diabetics. Stomach remedies, laxatives and diuretics may bring about mental changes through electrolyte disturbances and other mechanisms.

Physical examination is rarely helpful if the patient is "well" but localized neurological signs should obviously raise suspicions of a space-occupying lesion. The association of incontinence and ataxia or gait apraxia with mental disturbance may suggest normal pressure hydrocephalus, but there is considerable overlap with cerebrovascular disease⁵¹. Similarly, signs of Parkinsonism may be observed in patients with multi-infarct disease or diffuse Lewy body disease, who are unlikely to benefit much from L-dopa. On the other hand, patients with the characteristic extended posture and vertical gaze failure of progressive supranuclear palsy may respond to high doses of dopamine agonists⁵⁸. If the mental deterioration was preceded by a fall or there is evidence of head trauma, no matter how minor, suspicion of subdural haematoma should be aroused.

Some investigations, such as serum B₁₂ folate, calcium, thyroid function and screening for syphilis, are cheap to perform and lead to simple, if not always effective, medical treatment if an abnormality is found. It should be remembered that neuropsychiatric disorders caused by cobalamin deficiency⁵⁹ or folate deficiency^{60,61} can occur in the absence of haematological changes. On the other hand, the scientific evidence on which to base an effective and economically realistic policy for the use of expensive brain imaging techniques in people with dementia is still lacking.

Since the "treatable syndrome" of normal pressure hydrocephalus was first described by Adams *et al.*⁶² in 1965, there has been little agreement about the precise definition of the condition or the indications for treatment. Ventricular enlargement, with or without periventricular leukoaraiosis and various degrees of cortical atrophy, is a common CT finding in mentally normal old people, as well as those with dementia and other neuropsychiatric abnormalities. Clearly, it is not practical to monitor CSF pressure or to perform a lumbar infusion test in all cases, especially as such investigations do not always predict the response to internal shunting⁶³. Common sense would suggest that the response to repeated removal of CSF by lumbar puncture might be the best predictor of long-term benefit from surgery, but there again no systematic prospective study, let alone a randomized controlled trial, seems to have been done.

Similar, although less profound, uncertainty surrounds the treatment of subdural haematoma in the elderly. Here at least the CT scan should show a definite abnormality, but the relative merits of surgical and medical treatment (or none at all) are not precisely known. At the time of writing it is not possible to make logical recommendations for the clinical assessment and investigation of elderly people with dementia, since the issues are still surrounded by a fog of confusion and prejudice. The costs of investigation for all may be high, but they must be weighed against the enormous costs of institutional care and the psychological, social and indirect costs of the burden on carers, a substantial part of which might be avoidable. Large-scale, pragmatic outcome trials are urgently needed.

Depression and Functional Illness

In operational terms, if not according to strict definition, the commonest of the reversible dementias is so-called "depressive

pseudo-dementia". Here the psychiatrist has more to offer, but the physician is often involved because of the frequency of somatic symptoms and physical disabilities manifested by elderly depressed people. Indeed, when neither physicians nor psychiatrists are involved, people with potentially treatable illness can become heavy consumers of social services⁶⁴ or even residential care⁶⁵. In this situation the psychiatric history and mental state examination are of paramount importance, although yet again the possibility of adverse drug effects must be borne in mind. β -Blockers, methyl dopa and benzodiazepine are often implicated⁶⁶. Alcohol may be a cause or contributory factor and a simple screening questionnaire, such as the CAGE, should be included in any medical or psychiatric assessment⁶⁷. Physical examination is likely to be less rewarding. Although depression is a common complication of many physical illnesses, the proportion of cases where it is the sole presenting feature is quite small. Nevertheless, the chances of finding physical disease in elderly depressed patients are far higher than in their younger counterparts and its presence has a substantial adverse effect on prognosis: indeed, severe intractable depression in old age is nearly always associated with chronic physical ill-health⁶⁸. Secondary complications of depression, such as dehydration, nutritional deficiencies and constipation, must be identified and treated. It is unlikely that metabolic disturbances such as diabetes or hypothyroidism will be diagnosed clinically, so the appropriate blood tests should be done in all cases. A high ESR or C-reactive protein level should warn of the possibility of tuberculosis, other infection or cancer. An intensive search for occult neoplasia is rarely justified, although a chest X-ray should be done in any patient whose symptoms do not respond rapidly to treatment. Once again, the role of CT scanning will remain unclear until prospective studies of representative groups of old people presenting with clearly defined psychosyndromes have been reported. Finally, it should be remembered that the goal of clinical assessment in elderly patients is rarely to make a single unifying diagnosis. Multiple pathology is the rule rather than the exception and unusual combinations of symptoms or signs are more likely to be due to the combined effects of several common diseases rather than one rare one. The reduced homeostatic reserves of old age also means that a single initial insult (or treatment) often begins a chain of metabolic disturbances that multiply, leading to a cascade of complications. Thus, whatever the primary event, the clinician is often faced with a range of problems as well as pathophysiological diagnoses. Mental disturbance may therefore be seen as just one aspect of a complex multi-system disorder, but few would dispute that it is one of the most interesting and challenging of all.

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Needs and Problems

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Reliable and valid assessment techniques are needed for precision in studying, and communicating about, mental health problems. "Assessment" in this context conventionally refers to questionnaires and performance tests and behavioural observations can also be a part of a mental assessment¹⁻⁷.

The content of assessment is usually derived from a clinical experience⁸. However, clinicians may differ from each other, or from one interview to another, in the manner of their interview and their interpretation of the patient's responses. Therefore, reliability in assessment is obtained, in part, by prescribing the order and form of questions and tests (structuring the stimulus) and defining the informative response to be rated (structuring the response)⁹. This structuring of the interview technique enhances reliability, while retaining the focus of a clinically administered interview.

Structuring of mental assessment comes into its own when the condition has to be described in fairly subtle or phenomenological or behavioural terms, or when outcomes are influenced only to a limited degree by treatment or other factors, or where alternative treatments are not obviously different in their benefit and costs. Dramatic outcomes involving, for example, the difference between life and death, or the full recovery of an unusually incurable condition, hardly require precise measurements.

In an effort to reach precision in assessment through structuring of interview, construct validity may suffer. Conditions that are the focus of clinical concern but that cannot be precisely measured by state-of-the-art techniques may be excluded from research attention. Researchers may also choose to select types of patients not posing a problem for measurement. Alternatively, the concept of the condition may be constricted so that it fits within a measurable entity¹⁰. As the scope of psychiatric treatment expands, assessment techniques may have to follow suit.

A large proportion of elderly patients in institutional settings, and of patients with conditions such as dementia, or long-stay ageing schizophrenics in institutions, are unable to communicate at even a simple level. Testing or performance on cognitive tasks may not be possible, and questioning about inner feelings and thought processes is often completely out of reach. Attempts have been made to address these assessment problems by resort to informants^{11,12}, observations of reactions to non-verbal stimuli¹³ and inferences from responses to fairly straightforward stimuli, such as greeting behaviour. These approaches have not yet been proved to be entirely satisfactory.

The high correlation between mental and physical problems of the elderly makes it inevitable that mental assessment will often be confounded by physical symptoms. Somatic symptoms due to physical illness may register on scales intended to pick up the somatic manifestations of depression. Functional impairment due

to physical infirmity^{14,15} may spuriously fit criteria for determining the severity of dementia.

Various solutions to the confounding of the mental and physical symptoms have been proposed. Some investigators have chosen to develop instruments that concentrate on the mood state in depression^{16,17} or the cognitive state in dementia¹⁸, avoiding the need to attribute somatic or functional symptoms to a mental or physical cause. Other investigators have focused attention on the characteristics of somatic and functional symptoms that are suggestive of a mental rather than physical condition. Nevertheless, the measurement of the physical aspects of mental illness in the elderly still lacks specificity.

Potential users of mental assessment techniques are always concerned about the length of the time taken to administer the interview. Shorter interviews¹⁹⁻²² are viewed as more acceptable to subjects and less likely to be prematurely terminated, leading to a higher response rate for repeat interviews, cheaper to administer and less prone to error. There is no firm evidence that any of these assertions are true. Brevity has been carried perhaps to an extreme in assessments that serve as screening tools in busy clinical sites. Moreover, little attention has been given to devising interviews that are engrossing, reassuring and even helpful to the subject, so that longer interviews would be welcome.

Lengthening the time available for completing an interview allows more information to be collected on mental health and its associations. For most purposes in research and in clinical applications, there appears never to be enough time for all that might be relevant to study mental illness or clinical management. Brief scales of single dimensions (e.g. depressed mood or cognitive impairment) generally consume up to 10 min of interview time. Inclusion of material bearing upon differential diagnosis²³⁻²⁵ may extend to 30-40 min for current status and another 30 min or more for history of the illness. A comprehensive assessment^{9,25}, which covers the possible determinants and consequences of mental illness, may itself require 1 h or more, without exhaustive inquiry into any domain.

Compromises are usually necessary to keep the interview within feasible time limits while maintaining an adequate range of information. In this direction, contingencies may be introduced into the instructions for administration of the interview. Specified areas are probed only if a header question reveals the likelihood of useful information emerging from that area²⁶. There is a remarkable absence of empirical data showing that the rationale for these contingencies is well based. Furthermore, the complexity of contingencies may trap the interviewer to errors. Nevertheless, these contingent decision trees do represent the best chance of replicating the efficiencies and productivity of the way an experienced clinician conducts an interview. The advent of

computer-assisted technology for governing the conduct of interviews should facilitate the management of elaborate contingencies for driving the interview.

A certain line of questioning or testing may hinge upon a large body of information already gathered, rather than a single item or short set of items. A classic example is the use of an interview as a screen to select subjects for further investigation. Typically the latter investigation is at a separate time and venue, but it could follow straight on the heels of the screening interview at the same session. In either event it is essential to have immediate analysis of the information determining the contingency. Simple additive scoring systems are mainly invoked in these circumstances. Computer methods now allow much more elaborate analyses to determine the flow of sequential interviews or even sequences within a single session interview.

The capacity for rapid retrieval of assessment information on demand greatly enhances the value of mental assessment for supporting clinical decisions. Yet the potential in this respect cannot be fully realized until clinical goals of assessment assume priority over research goals in shaping the form and content of the assessment. Notwithstanding that mental assessment emerged from clinical experience, the current state or nature of mental assessment owes much more to research than clinical interests. A return to the origins of its development is called for, in order to bring assessment into line with clinical as well as research needs^{27,28}.

Clinical activities, for reasons of practice organization and fiscal considerations, generally take place under pressure of time. The collection and even analysis of information for the review clinician can be delegated to personnel under less time pressure than the physician/psychiatrist. In that arrangement, it becomes essential that the transfer of crucial information to the physician be managed expeditiously. This entails highly discriminating summarization, display and communication of mental assessment information. It also assumes that the physician learns to assimilate the type and form of information and to incorporate it in the planning and monitoring of clinical management. This learning process, like other clinical skills, must be continuously honed through experience assisted by consultation. There are few guidelines as yet as to the assignment, conduct and necessary training for the professional role of marshalling the mental assessment information and communicating it to clinicians.

Much of mental assessment is aimed at documenting the subjective experience or inner states of a patient²⁹. The best reporter for this material should be the patient, but the latter may be uncommunicative or give misleading information. An informant may not do much better. Under these circumstances, it would be desirable to turn to objective testing and observation of behaviour³⁰⁻³² and laboratory results if these can be obtained. As matters stand, laboratory findings in themselves are at best ambiguous³³ as mental assessment information in the elderly, although there is hope that eventually laboratory tests will play a larger part in mental assessment. In any event, whether intended or not, many subjects in epidemiological studies will not undergo laboratory testing. In the meantime, other ways of achieving objectivity are more likely to be productive. Testing organic mental status is objective in the sense that the task performance of the person is observable and errors can be quantified. Programming is the presentation of the tests through computer software and adds to the objectivity^{13,34}. Testing of an objective nature has also evolved for functional performance for tasks involving everyday activities, or simulations of the skill demands entailed in such activities^{14,15}. Inefficiencies in carrying out these tasks provide information on the diagnosis of certain mental conditions (e.g. dementia) and the outcomes of a wide range of mental illnesses.

Structuring of testing procedures (backed up by suitable training) allows the results of assessment to be compared with

data collected on the same instruments in different studies. It also permits data to be interpreted in the light of accumulated experience on the normal distribution and longitudinal course of levels or patterns of test results in the general population and in specific clinically defined groups. Thus, the value of assessment information is enhanced in diagnoses, prediction of outcomes, selection of treatment and evaluating the significance of changes in levels or patterns of test scores. This full psychometric development warrants the use of the term "standardized" to describe a method (although the term is used to mean structured). However, there are several cautions that must be addressed in attempting to draw upon the potential value of standardization.

Unless an assessment technique is administered in identical fashion in two or more studies, the results of those studies cannot be directly compared and information cannot be transferred from one study to another. Not only must the structure and procedures of the assessment protocol be kept constant, but also the methods of training. Furthermore, adjustments must be made for differences in the characteristics (e.g. age, education and culture) of two populations that might alter the meaning and confound the comparison of results from the same assessment technique³⁵. These adjustments may be difficult to make unless standardization has been accomplished on appropriate populations and formulae for interpreting scores have been worked out^{36,37}. Attempts are being made to construct culture-free assessments which, in effect, would obviate the need to adjust scores with reference to demographic characteristics.

Most assessment techniques devoted to the classification of mental disorders still rely heavily upon information gathered at one point in time³⁸, cross-sectional status with or without retrospective historical information. Longitudinal or prospective information is principally regarded as measuring change, course and outcomes. Yet the longitudinal picture of mental health problems offers crucial clues to diagnosis and needs to be incorporated into assessment techniques for classification. In order to meet the demands of longitudinal measurements, assessment techniques must deal with practice effects, ceiling and floor limits to the range of measurement, and the minimization of attrition by maintaining the interest of the subject.

Certain domains of mental ill-health appear well-represented, even over-represented, in current assessment techniques. For instance, there are probably more measures of cognitive impairment^{19,25,39-47} and of depressed mood^{16,17,48-56} than are strictly needed. Conversely, there are neglected domains. Innovations would be welcome in the measurement of positive mental health, either as a global concept or as applied to specific mental health areas, such as affect⁵⁷ and cognition. There is a need to understand the contribution of the measurements of positive states to diagnosis and prognosis.

More generally, the scope of assessment techniques should be expanded and explored so as to fill in gaps in the domains relevant to describing mental health and its associations. Moreover, assessments need to encompass the various perspectives of subject, family member^{11,12,58,59}, health professional and other parties with a legitimate interest in the subject's mental health. When a complete inventory of domains and perspectives has been captured by assessment techniques, it may become possible to describe a reasonable approximation to the plight of the whole person.

A note on quality of life assessment. The concept and measurement of quality of life has come to fill a prominent place in geriatric health care. Since about 1970 the relevant professional literature on measurement and its applications has steadily increased in volume. Instruments have proliferated and some have assumed ascendancy in terms of widespread use. There is no better indication of the need for measurement of this concept than the many applications to which it has been turned. These

include descriptions of the course of illness, comparisons of the outcomes of competing treatments, monitoring of the performance of healthcare systems, accounting for benefits derived from resources allocated to various interventions, assisting decisions on treatment choice, and informing policy formation on development of the health care system and its components.

Yet there are problems of concept and measurement that have emerged for which no satisfactory solutions have yet been found. For example, there are unresolved conflicting advantages and limitations offered by generic or specific measures, by emphases on subjective or objective approaches, by brief or comprehensive assessments, or by combining or segregating status and preferences.

Many of the existing quality of life instruments have adopted scales and indices that had a prior existence as measures of health status. This has created a core of domains that are found in a majority of instruments: this core includes functioning in the activities of daily living, mobility, cognitive status, depression or morale, physical discomfort and self-perceived health. However, there are many other domains that appear in some instruments and not in all, so that the potential cumulative list is lengthy. There is little empirical work to identify which domains are most critical to a good quality of life.

Choosing between the numerous established instruments is a daunting task for the relatively inexperienced researcher or for the clinician wishing to enhance practice standards. An instrument which is widely used may appeal to researchers, reviewers and grant committees on the grounds of its substantial psychometric development, provision of norms, comparison with findings from previous pertinent studies, as well as the credence that comes from the consensus implicit in a large constituency of users.

Nevertheless, there are junctures in the growth of a field within the health sciences at which technological expediency can outstrip the slower work of fundamental understanding. At such points in the history of the maturation of a field the good can pre-empt the excellent. Whether this is the case for the current state of quality of life concept and measurement is a matter of judgment.

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Non-computerized Assessment Procedures: Fundamental Assessment Issues

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BASIC ASSESSMENT ISSUES

Just as individuals age, so do populations. The “graying of America” is a cliché that carries with it the inevitability of the normal aging process combining with the advancement of medical science. The average age of all industrialized countries is going up steadily. While, to some extent, these changes in basic demographics relate to improved infant mortality, there is also an increasing ability and willingness on the part of the medical profession to treat the ailments that may come with aging and extend the lifespan of individuals within the society. Control of ischemia risk factors, dialysis and organ transplant are all commonplace now but were much less advanced 30 years ago.

In many ways this demographic change is a very positive trend. Individuals at 50 now are no longer considered to be as old as they once were. There are many physically vigorous and cognitively intact individuals who function effectively well into their 70s, 80s and 90s. However, extension of life is not necessarily the extension of quality of life. With increasing age there is an increasing probability of dementing processes. Dementia is not the inevitable outcome of aging, but the diagnosis of dementing conditions increases significantly with each decade of life.

There are multiple potential dementing conditions. Some of these conditions are the by-products of physical disease and others of lifestyle choices made years earlier (*see* Table 27.1 for a partial listing of the more common dementing processes from the familiar DSM). The clear majority of dementing processes, however, fall into three general categories, Alzheimer’s dementia, vascular disease-related dementia, and mixed dementias. While each has a progressive course, the potential for treatment and the speed and uniformity of the progression varies significantly. Precision of diagnosis is critical in the appropriate treatment and management in a geriatric dementia work-up.

Depression in the old, in combination with the normal aging process, can produce a condition that is treatable but that mimics the effects of central nervous system-based dementias. Depression in some cases may be the earlier preclinical precursor of dementia, but nevertheless, the accurate differential diagnosis of dementia and depression is of extraordinary importance. The consequences of a false-positive or false-negative diagnosis in dementia can be devastating. If a patient is labeled as having early-stage Alzheimer’s syndrome, multiple negative consequences occur very quickly. Driving privileges, financial and medical decision-making and the independence that is critical to any person are all restricted and sometimes taken away. If the patient diagnosed with dementia is not demented but is depressed, unnecessary and

Table 27.1 DSM-IV: common dementing processes

Dementia of the Alzheimer’s type
Vascular dementia
Dementia due to HIV disease
Dementia due to head trauma
Dementia due to Parkinson’s disease
Dementia due to Huntington’s disease
Dementia due to Pick’s disease
Dementia due to Creutzfeldt–Jakob disease
Dementia due to other general medical conditions
Substance-induced persisting dementia
Dementia due to multiple etiologies
Dementia not otherwise specified

Diagnostic and Statistical Manual of Mental Disorders, 4th edn, p. 133.

potentially irreversible changes occur in the quality of the patient’s life, even if the diagnosis is later changed. Similarly, if the patient does have a progressive dementia and proceeds to make decisions about finances, about health, or even about day-to-day instrumental activities of daily living, the consequences can be equally negative.

In summary, the geriatric population is growing. As people age, they become more susceptible to potentially treatable conditions. These conditions can radically alter the quality of life and the ultimate prognosis and disposition for the successful management of the case.

ASSESSMENT PROCEDURES

The assessment of geriatric patients for dementing conditions demands considerable focus in assessment and knowledge of the scientific literature. These areas would include, at minimum, an understanding of the characteristics of the normal aging process and the salient behavioral/cognitive characteristics of the dementias. Necessary and desirable parameters of geriatric evaluations are as follows.

Necessary Characteristics of Any Assessment Procedure

Norms

Test results are significantly influenced by the age of the subject and premorbid functioning levels; therefore, any assessment

procedure utilized should have reasonably adequate norms that take age and educational level into consideration.

Validity and Reliability

Validity and reliability of the assessment procedure must be established and the relationship between the given dementing process and test scores well established within the scientific literature.

Depression and Cognition

Cognitive and affective diagnostic procedure should be utilized within the same battery, whether formally or informally, since the two interact so powerfully. The diagnosis of dementia vs. depression often resolves itself down to the proportion of each.

Depth|Breadth of Assessment

Sufficient breadth and depth of assessment is required so that patients can be followed over time, in improving and deteriorating conditions, so that unsuspected conditions are not easily missed in a routine assessment.

Desirable Characteristics

Short

Relative brevity, even in a multi-factorial assessment, is highly desirable. Geriatric patients become impatient, threatened and exhausted when test batteries become too lengthy. The procedure at the end of the assessment process may be less valid than that at the beginning of the assessment process.

Practical

Ideally, along with reliability and validity, the assessment encompasses those variables that are clinically important. While reaction time may be highly sensitive at the bedside, in day-to-day clinical practice memory assessment or assessment of attention is much more likely to be broadly useful. The geriatric patient is more likely to be uncooperative and sometimes appropriately contemptuous of “kid” games. If a patient can perceive the relevance of the procedure, whether or not he/she feels threatened by it, the test results are more likely to be valid and reliable.

Numbers

Quantification of the test result is very desirable, since multiple successive assessments may be done that trace the course of the syndrome or the effects of medication.

Models

It is highly desirable that the tests, in aggregate, fit together into a relatively coherent model of cognitive assessment and include formal and informal assessment of at least the following:

- General abilities.
- Speech and language.

- Constructional abilities.
- Motor functions.
- Memory and learning.
- Attention and concentration.
- Judgment and problem-solving abilities.
- Speed of processing.

At times, these assessments may be as informal as just listening to the patient, but are necessary descriptive parameters of the assessment.

NON-COMPUTERIZED ASSESSMENT OF COGNITIVE FUNCTION

Selection of Assessment Tools

As in all patient populations, neuropsychological assessment of older individuals begins with a thorough understanding of the referral question and a clear goal as to the purposes of the evaluation. The specific functional areas of cognition to be tested, and therefore the format and comprehensiveness of the entire evaluation, should be based on the initial referral question and targeted at achieving the specific goals of the referring healthcare provider. Nowhere is it more important to tailor the assessment battery to the specific diagnostic concerns, while considering the unique abilities and limitations of the patient population, than with a geriatric population. Test selection must optimally allow for an in-depth evaluation of the cognitive areas of concern, while keeping battery length and difficulty level manageable for the aging patient. Ultimately, the battery “should discriminate maximally between normal aging and CNS disorders such as the different dementias”¹. As these authors state further, a successful test battery should also allow differentiation among the various common subtypes of dementia, as well as between dementia and affective disturbance.

Screening Evaluations vs. Full Batteries

Even when the referral question implies a broad assessment of overall cognitive abilities for the purpose of identifying the presence or absence of neuropsychological deficits, the use of a brief but well-rounded neuropsychological screening measure can be prudent. Such tools typically allow the examiner to quantify gross deficits against normative data in order to classify the tested individual as “normal” or “abnormal” in particular cognitive areas of functioning. Cut-off scores and the number of abnormal scores necessary for rating an individual’s performance on such measures against target normative populations usually allow for qualitative classifications ranging from superior to severely impaired. Depending upon the psychometric soundness of the screening instrument, the examiner may be able to draw conclusions with respect to specific cognitive abilities, or may be limited to a judgment of impaired or not impaired.

The benefits of using a brief screening measure for cognitive evaluation are obvious; they tend to require less administration time for both the patient and the examiner, and they are typically less labor-intensive and intimidating for the patient. Most commercially available cognitive screens are easily scored and provide feedback for recommendations quickly, which is an asset in inpatient settings or other situations when the patient’s treatment plan and disposition considerations may be urgent. As with most things in life, however, time-saving procedures may sacrifice quality. Even the most widely accepted brief cognitive screens lack the diagnostic sensitivity and specificity of the more comprehensive neuropsychological battery, particularly when

they generate only a single score or are limited to largely verbal measures. When choosing to use a cognitive screen for neuropsychological evaluation of a geriatric patient, it is important to balance our desire to spare the patient a lengthy assessment time with a genuine quest to garner the most reliable diagnostic data available. This goal may best be achieved by augmenting a brief cognitive screen, with individual neuropsychological measures aimed at the more detailed evaluation of specific cognitive abilities or affective characteristics.

One neuropsychological screening assessment that has achieved wide acceptance with use in a geriatric population is the Dementia Rating Scale (DRS)². This scale yields scores for attention, initiation/perseveration, construction, conceptualization and memory, and compares individuals' performances to a normative base of subjects with known Alzheimer's disease. In providing such a comparison population, the instrument controls for normal aging variation. In this way, the measure offers easily referenced cut-off scores for quick classification of outcomes as being consistent with dementia or not.

A more recent tool available to clinicians is the Repeatable Battery for the Assessment of Neuropsychological Status (RBANS)³. While supportive literature for its use in diagnosing dementia in the geriatric population is only now being gathered, the RBANS has the advantage of being available in parallel forms, thus allowing serial assessment. Repeatability is a valuable feature for a cognitive screen to have, especially in documenting recovery of function after a reversible neurological injury and in demonstrating progressive decline in the degenerative dementias. The authors refer readers to published review articles and commercial catalogs for the clinical findings and availability of these and other neuropsychological screening measures.

In many cases, the discrimination between dementia and pseudodementia, or the differential diagnosis between types of dementing illnesses, cannot be achieved through use of a brief cognitive screen. In these cases, a more comprehensive test battery may be the only method by which to quantify and qualify specific areas of deficits into a pattern consistent with a diagnostic picture. In selecting tests for such a battery, one may focus on specific symptomatic cognitive areas or may seek to assess global cognitive functioning with a combination of neuropsychological measures. There are numerous individual neuropsychological tests developed to assess specific cognitive functional areas (Table 27.2). The combination of these tests into a clinical battery may be based on theoretical concepts or assessment approaches; some clinicians prescribe to use of standard batteries, such as the Halstead-Reitan and the Luria-Nebraska, while others combine individual neuropsychological measures into a more flexible or "process approach" battery. Whatever one's theoretical basis, consistency and rigor in assessment procedures remains the most effective way for a clinician to develop his/her own personal bank of base rates and characteristic result profiles.

Specific Cognitive Functional Areas

Intellectual functioning is one of the broader functional areas assessed and may include assessment of fluid and crystallized knowledge. It is not a unitary entity but instead consists of multiple functions, including the ability to acquire, process, categorize and integrate information. Intellectual functioning intimately involves use of memory and learning, visuospatial skills, attentional abilities, expressive and receptive language, and aspects of adaptive reasoning and organizational structure. Assessment may involve evaluation of verbal intellect, non-verbal (performance) intellect, or overall intellectual abilities.

Attention and concentration abilities are critical for neuropsychological functioning. More complex processing depends on

Table 27.2 Examples of neuropsychological tests for specific cognitive functions

Cognitive functional area	Example neuropsychological tests
Intellectual functioning	WAIS-R or WAIS-III Ravens Progressive Matrices (non-verbal) Peabody Picture Vocabulary Test (verbal) Mini-Mental State Examination (MMSE)
Attention/concentration functioning	Verbal Series Attention Test Continuous Performance (2 and 7) Test Stroop Test Digit Span PASAT
Executive functioning	Wisconsin Card Sorting Test Short Category Test Trail Making Test Verbal Fluency/Figural Fluency
Memory and learning functioning	WMS-R or WMS-III (logical memory and visual reproduction) CVLT, RAVLT-R, HLVLT-R Selective Reminding (Buschke or Levin) Warrington Facial Recognition Memory
Language functioning	Benton Visual Retention Test Western Aphasia Battery Wepman Aphasia Screening Multilingual Aphasia Examination Boston Naming Test Controlled Verbal Fluency Animal Fluency
Visuospatial/visuomotor functioning	Benton Facial Recognition
Judgment of line orientation	Rey-O Complex Figure Trail Making Test
Motor functioning	Finger Tapping Grooved Pegboard Grip Strength Boston Apraxia Examination
Emotional/personality functioning	Geriatric Depression Scale Beck Depression Inventory Multiscore Depression Inventory MMPI-2

these primary functions. Their scope involves speed of information processing, accuracy of discrimination, selectivity and use of a heightened state of awareness. Successful evaluation of attention/concentration skills is necessary for proper interpretation of other neuropsychological findings, including memory and learning.

Executive functioning is a complex phenomenon, which answers the question of how or whether one will do something. It involves the capacity to initiate activity and the process of self-monitoring and interpreting feedback. Executive functions include the planning and sequencing strategies that facilitate goal-directed behaviors. Assessment of executive functioning should include testing at varying levels of complexity to obtain an adequate picture of the patient's mental flexibility.

Memory and learning assessment is at the core of most neuropsychological evaluations of the elderly. A complete evaluation should examine verbal memory (contextual and non-contextual) and visual memory (independent of constructional ability), and should address acquisition, retention and recognition through somewhat independent means. The distinction between deficits in encoding and deficits in retrieval of newly-learned information can be crucial in differential diagnosis of the various dementias.

Expressive and receptive language abilities should be assessed directly, including auditory and reading comprehension, repetition skills and the ability to follow simple commands (praxis). Specific language characteristics, including visual confrontation

naming and word fluency, may be evaluated with multiple or overlapping measures to increase the reliability of findings. Whenever possible it may be helpful to assess language abilities in both the oral and written modalities to determine the generalizability of language deficits.

Visuospatial functioning assessment should include the evaluation of near visual acuity, processing of visual stimuli, recognition of shapes, colors and forms, and overall spatial orientation of objects in space and during motor tasks. The copying and drawing of geometric or figural designs can directly assess visual perception.

Assessment of motor functioning should include both gross and fine motor abilities. Manual dexterity and gross manual strength are both important and may yield clues to hemispheric differences. Motor planning can best be evaluated through completion of complex, repetitive motor movements and with traditional go–no go motor tasks.

Emotional/personality functioning is one of the most important areas to be assessed in the elderly population. Affective disturbance can contribute to multiple cognitive deficits; therefore, the differentiation between emotional dysfunction and true CNS abnormality is essential in creating an accurate diagnostic profile. Generational differences must be considered when assessing elderly patients for signs and symptoms of depression and anxiety, and stereotypical views of expected emotional functioning of older individuals must be avoided.

ADDITIONAL RESOURCES

What has been offered in this chapter represents a brief introduction and overview of the basic concepts of non-computerized neuropsychological assessment in geriatric patients. The field of geriatric neuropsychology, while still relatively new in comparison with other disciplines, is growing exponentially as the world population continues to include more people at older ages. There exist a good number of valuable references and resources

for both professionals working with older individuals, and for caregivers who must consider whether referral of their loved one for neuropsychological evaluation is indicated. The authors encourage healthcare professionals and caregivers alike to access their local centers on aging, the national Alzheimer's Association, and the various clinical neuropsychological associations (e.g. National Academy of Neuropsychology) to obtain further information on cognitive assessment and intervention in older individuals. An extensive list of reading references is also being included to encourage further exploration of related topics.

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Mini-Mental State Examination

Joseph R. Cockrell and Marshal F. Folstein*

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The Mini-Mental State Examination (MMSE) is a 10-minute bedside measure of impaired thinking in undeveloped, uneducated, diseased, or very old populations. The summed score of the individual items indicates the current severity of cognitive impairment. Deterioration in cognition is indicated by decreasing scores of repeated tests. Scores are reliable between tests and between raters¹ and correlate with other mental tests, electroencephalography², computerized tomography³, magnetic resonance imaging⁴, single photon emission computed tomography (SPECT) scan⁵, CSF proteins and enzymes^{6,7} and brain biopsy synapse numbers⁸.

To administer the MMSE, gain the patient's cooperation by asking him if there is a memory problem and then asking permission to test his memory. During testing, praise successes and ignore failures so as to avoid emotional reactions that will compromise cooperation and performance.

The items of the MMSE include tests of orientation, registration, recall, calculation and attention, naming, repetition, comprehension, reading, writing and drawing. Level of consciousness is rated on a scale from coma to fully alert, but the consciousness rating is not summed with the other items. If all items are answered correctly, the score is 30. The mean score for a community-dwelling population over 65 years of age is 27, with a standard deviation of 1.7¹. The score is lower in those who completed comparatively fewer years of education and who have diagnosable diseases^{9–11}. Patients with dementia, delirium, mental retardation, Parkinson's disease, stroke and some cases of depression score lower than normal controls^{1,12}. Alzheimer's disease patients lose 3–4 points per year of illness after the onset of memory disturbance, although there is wide variability in this phenomenon^{13,14}.

naming and word fluency, may be evaluated with multiple or overlapping measures to increase the reliability of findings. Whenever possible it may be helpful to assess language abilities in both the oral and written modalities to determine the generalizability of language deficits.

Visuospatial functioning assessment should include the evaluation of near visual acuity, processing of visual stimuli, recognition of shapes, colors and forms, and overall spatial orientation of objects in space and during motor tasks. The copying and drawing of geometric or figural designs can directly assess visual perception.

Assessment of motor functioning should include both gross and fine motor abilities. Manual dexterity and gross manual strength are both important and may yield clues to hemispheric differences. Motor planning can best be evaluated through completion of complex, repetitive motor movements and with traditional go–no go motor tasks.

Emotional/personality functioning is one of the most important areas to be assessed in the elderly population. Affective disturbance can contribute to multiple cognitive deficits; therefore, the differentiation between emotional dysfunction and true CNS abnormality is essential in creating an accurate diagnostic profile. Generational differences must be considered when assessing elderly patients for signs and symptoms of depression and anxiety, and stereotypical views of expected emotional functioning of older individuals must be avoided.

ADDITIONAL RESOURCES

What has been offered in this chapter represents a brief introduction and overview of the basic concepts of non-computerized neuropsychological assessment in geriatric patients. The field of geriatric neuropsychology, while still relatively new in comparison with other disciplines, is growing exponentially as the world population continues to include more people at older ages. There exist a good number of valuable references and resources

for both professionals working with older individuals, and for caregivers who must consider whether referral of their loved one for neuropsychological evaluation is indicated. The authors encourage healthcare professionals and caregivers alike to access their local centers on aging, the national Alzheimer's Association, and the various clinical neuropsychological associations (e.g. National Academy of Neuropsychology) to obtain further information on cognitive assessment and intervention in older individuals. An extensive list of reading references is also being included to encourage further exploration of related topics.

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Misinterpretations of scores result from several misconceptions about the test. The MMSE is not a complete mental status examination or a complete neuropsychological examination. The MMSE does not define a clinical or pathological diagnostic category, such as dementia or brain tumor or organicity. The score does not measure decline from a previous level unless tests are repeated over time. The score does not tell the whole story. Individual items are useful for understanding the situation of the patient, since they indicate whether the patient can follow instructions, read and write. Finally, the MMSE is a weak measure of competence or disability. Competence, handicap and disability must be assessed by procedures designed for that purpose.

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IQCODE: Informant Interviews

A. F. Jorm

Centre for Mental Health Research, The Australian National University, Canberra, Australia

In assessing patients for dementia, informants are a valuable source of information, complementing other sources such as cognitive testing. The strengths of informant data include:

- *Everyday relevance.* Informants can report on how the patient is functioning in everyday cognitive tasks. Cognitive tests, by contrast, generally involve artificial tasks removed from daily life.
- *Acceptability to patients.* Formal cognitive testing can be distressing to some people because of the limitations it reveals. However, informant interviews do not directly confront the patient's limitations.
- *Longitudinal perspective.* It is often useful to know how a patient is functioning compared to earlier in life. An interview with an informant who has known the patient for many years can provide this.
- *Ease of administration.* If necessary, informant data can be collected by telephone or mail. In some research situations, it has even been used to assess deceased subjects.
- *Cross-cultural portability.* Informant data may have greater validity than cognitive testing for patients who are from culturally different backgrounds¹.

While informant interviews are widely used in clinical practice, there is now a range of standardized informant interviews and

questionnaires available for quantifying this information. Probably the most widely used and researched is the Informant Questionnaire on Cognitive Decline in the Elderly (IQCODE)². The IQCODE is a 26-item questionnaire that asks the informant about cognitive changes over the previous 10 years. Items cover memory and intellectual functioning and are rated on a five-point scale from “1. Much improved” to “5. Much worse”. Originally, the IQCODE was designed as an interview but it is more commonly used as a self-administered questionnaire. There is now a short 16-item IQCODE which performs as well as the original. Translations of the IQCODE are available in a range of languages. The various versions are available on the Web at <http://www.anu.edu.au/iqcode/>.

Principal component analysis of the IQCODE items have shown that it measures a general factor of cognitive decline. Validity studies have found that the IQCODE correlates moderately with cognitive screening tests such as the Mini-Mental State Examination (MMSE) (mean $r=0.59$ over seven samples). Correlations with indicators of premorbid ability, such as the National Adult Reading Test (NART) and years of education are repeatedly found to be near zero.

A meta-analysis of seven studies directly comparing the IQCODE with the MMSE as a screening test for dementia found that the IQCODE performed at least as well as the

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Staging Dementia

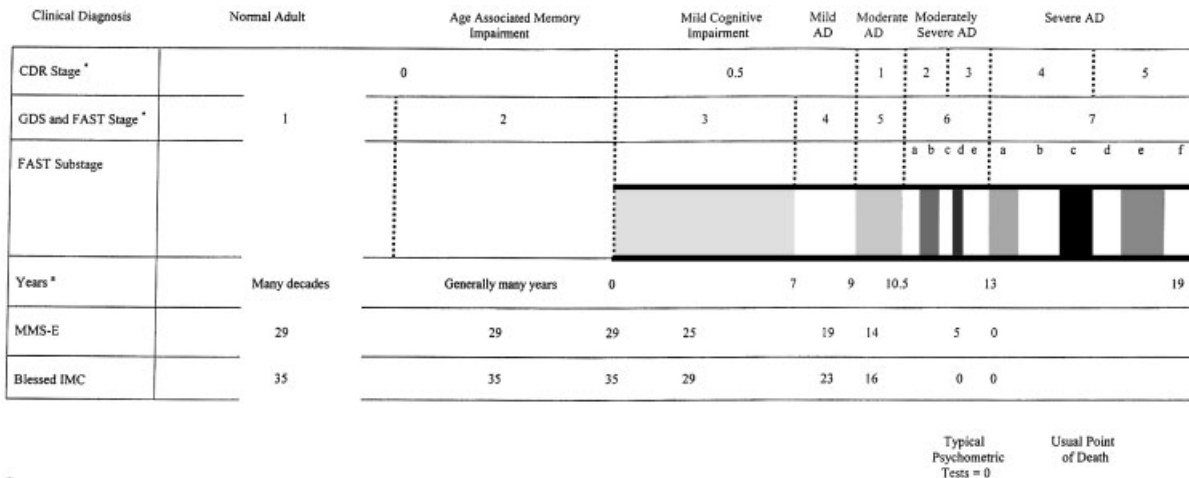
Barry Reisberg, Gaurav Gandotra, Arshad Zaidi and Steven H Ferris

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Dementia is a progressive pathologic process extending over a period of many years. Clinicians and scientists have long endeavored to describe the nature of this progression. Such descriptions have generally been encompassed within two broad categories, viz. global staging and more specific staging, sometimes referred to as axial or multi-axial staging. A comparison of the major current dementia staging systems with the most widespread mental status assessments in Alzheimer’s disease, the major cause of dementia, is shown in Table 1, which illustrates some of the major potential advantages of staging. These advantages include: (a) staging can identify premorbid but potentially manifest conditions that may be associated with dementia, such as age-associated memory impairment, a condition which is not differentiated with mental status or psychometric tests; (b) staging can be very useful in identifying subtle, identifiable, prodementia states, such as mild cognitive impairment (MCI), wherein mental status assessments and psychometric tests, while frequently altered, are generally within the normal range and consequently are not reliable markers; and (c) staging

can track the latter 50% of the potential time course of dementias such as AD, when mental status assessments are virtually invariably at bottom (zero) scores. Furthermore, apart from its utility in portions of dementia where mental status and psychometric assessments are out of range or clearly insensitive, there is evidence that staging procedures can more accurately and sensitively identify the course of dementia in the portion of the condition that is conventionally charted with mental status assessments. This latter evidence comes from longitudinal investigation of the course of AD¹, pharmacologic treatment investigation of AD² and study of independent psychometric assessments of AD³. Another seeming advantage of staging procedures in comparison with mental status or psychometric assessment of AD and other dementias is in identifying the management concomitants of severity assessments⁴. Staging procedures have also been successfully applied post mortem to assess retrospectively the diagnoses of a diverse assortment of dementia-related cases available for “brain banking” but on which no ante mortem clinical data were available⁵. Similarly,

Table SA27iii.1 Typical time course of normal brain aging and Alzheimer’s disease



*Stage range comparisons shown between the CDR and GDS/FAST stages are based upon published functioning and self-care descriptors.

^aNumerical values represent time from the earliest clinically manifest symptoms of Alzheimer’s disease.

Adapted by permission from Reisberg *et al.*⁵²

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Table SA27iii.1 Typical time course of normal brain aging and Alzheimer's disease

Clinical Diagnosis	Normal Adult	Age Associated Memory Impairment	Mild Cognitive Impairment	Mild AD	Moderate AD	Moderately Severe AD	Severe AD	
CDR Stage *		0	0.5	1	2	3	4	5
GDS and FAST Stage *	1	2	3	4	5	6	7	
FAST Substage					a	b c d e	a b c d e f	
Years *	Many decades	Generally many years	0	7	9	10.5	13	19
MMS-E	29	29	29	25	19	14	5	0
Blessed IMC	35	35	35	29	23	16	0	0

Typical Psychometric Tests = 0 Usual Point of Death

*Stage range comparisons shown between the CDR and GDS/FAST stages are based upon published functioning and self-care descriptors.

^aNumerical values represent time from the earliest clinically manifest symptoms of Alzheimer's disease.

Adapted by permission from Reisberg *et al.*⁵²

post mortem retrospective staging procedures have been successful in establishing remarkably robust clinicopathologic correlations in longitudinally-studied AD cohorts^{6,7}.

GLOBAL STAGING

Efforts to stage progressive dementia globally can be traced back at least to the early nineteenth century, when the English psychiatrist, James Prichard, described four stages in the progression of dementia: “(1) impairment of recent memory, (2) loss of reason, (3) incomprehension, (4) loss of instinctive action”⁸. More recently, the American Psychiatric Association’s 1980 *Diagnostic and Statistical Manual of Mental Disorders*, 3rd edn (DSM-III)⁹ recognized three broad stages in its definition of primary degenerative dementia. Subsequently, in 1982, two more detailed global descriptions of the progression of dementia were published. One of these, the Clinical Dementia Rating (CDR) scale¹⁰ described five broad stages from normality to severe dementia. The other, the Global Deterioration Scale (GDS)¹¹, identified seven clinically recognizable stages, from normality to most severe dementia of the Alzheimer type. These two global staging instruments, the GDS and the CDR, are generally compatible except that the GDS is more detailed and specific and identifies two stages that the original CDR staging does not. One of these is a stage in which subjective complaints of cognitive deficit occur (GDS stage 2). These subjective complaints are now recognized as occurring very commonly in aged persons¹²⁻¹⁴ and consensus workgroups have called attention to the importance of these symptoms and the need for more detailed study of their nature and treatment¹⁵⁻¹⁷. Although this stage of subjective complaints continues to be identified only by the GDS staging system, recent studies have indicated that persons with these complaints are at increased risk for subsequent overt dementia^{18,19}. Also, at the other end of the pathologic spectrum, the CDR did not identify any stage beyond that in which dementia patients “require much help with personal care” and are “often incontinent”, whereas the GDS identifies a final seventh GDS stage in which patients are already incontinent and over the course of which language and motor capacities are progressively lost. Subsequently, two further stages were suggested for the CDR, corresponding to the GDS stage 7 range²⁰⁻²².

Staging procedures have been shown to be valid and reliable methods for assessing the magnitude of pathology in AD and related dementing conditions. This validity and reliability is illustrated in this brief review for the GDS, probably the most detailed and explicit staging procedure.

The validity of the GDS has been demonstrated in several ways. Cross-sectional studies have confirmed the consistency of the ordinal sequence and the optimal weighting of the hierarchically sequenced items embodied in the GDS stages in aging and progressive Alzheimer’s disease (AD)²²⁻²⁴. Thus, the specific impairments characteristic of each stage almost always follow the impairments described for the previous stage. Also, the grouping of impairment characteristics within stages appears to be optimal.

For example, naturalistic study has supported the identification of staging phenomena, largely identical to the GDS stages, by independent lay-person observers. In this study²³, a 30-item questionnaire derived from the GDS was completed by a relative or caregiver for each of 115 patients with varying degrees of dementia. Principal components analysis was used to combine the items into a single composite scale which more reliably represents distances between the 30 clinical manifestations along the continuum of cognitive decline. The study found that “the scale scores for the clinical manifestations were observed to cluster into relatively discrete groups, suggesting naturally occurring stages or

phases. Objective cluster analysis methods further confirmed the presence of distinct transitions along the cognitive decline continuum”. It concluded that the “utility of empirically derived scale values in staging the course of primary degenerative dementia is suggested”.

The relationship between the GDS stages and mental status assessments, other dementia assessments, scores on cognitive tests and other objective tests and *in vivo* assessments of brain change in aging and progressive dementia have been studied in considerable detail^{11,24}. These studies have indicated significant correlations between all of these measures of dementia severity and the GDS stages. However, the strongest relationships have been observed between comprehensive dementia assessments, such as the Mini-Mental State Examination (MMSE)²⁵, and the progression of dementia on the GDS²⁴. The GDS also correlates well with evaluations of actual functioning and activities of daily living in AD²⁶ and with independent physical markers of AD progression, such as changes in neurologic reflexes²⁷. Thus, the construct validity of the GDS has been well substantiated.

At least six separate studies have examined the reliability of the GDS²⁸⁻³³. Reliability coefficients have ranged from 0.82 to 0.97 in these studies, using disparate procedures in diverse settings. These studies have indicated that the GDS is at least as reliable as any other instrument upon which clinicians rely, such as the MMSE. In a reliability study in a nursing home setting³², the GDS was found to be somewhat more reliable than the MMSE. Importantly, GDS staging has also been shown to be a reliable procedure when assessed using a telephone format³³.

Global staging scales such as the GDS have certain important advantages in dementia assessment. First and foremost, these scales are strongly anchored to the clinical symptoms, behaviour and functional changes in progressive degenerative dementia and particularly that of Alzheimer’s disease. Consequently, they discourage misdiagnosis. Unlike many mental status and other dementia test instruments, global stages are relatively stable over time and relatively resistant to practice effects. Equally importantly, global staging instruments are minimally influenced by educational background and socioeconomic status, whereas mental status and similar assessments are strongly influenced by such factors. Also, global staging, and in particular the GDS, covers the entire range of pathology in central nervous system (CNS) aging and progressive dementia, whereas, for example, mental status assessments and most psychometric tests entirely fail to distinguish GDS stages 1 and 2. Occasionally, patients may display GDS stage 3 symptomatology and still score a perfect 30 on the MMSE. Uncommonly, dementia patients may display GDS stage 4 symptomatology, and still score a perfect 30 on the MMSE. Much more commonly, patients may display the clear-cut dementia symptomatology characteristic of GDS stage 4 and achieve MMSE scores which are near-perfect or within the so-called “normal” range. At the other end of the pathologic spectrum, most patients at GDS stage 6 achieve only bottom scores on traditional psychometric tests. Over the entire course of the GDS 7 stage, nearly all patients attain only zero scores on the MMSE. The GDS, however, describes a final seventh stage, over the course of which patients may survive for many years.

AXIAL AND MULTI-AXIAL STAGING

The observation that the progression of dementia pathology is accompanied by progressive changes in more specifically defined processes has resulted in efforts to stage dementia in terms of those processes. Generally, axial staging has attempted to exploit progressive changes in cognition or functioning, although attempts have also been made to stage hierarchically progressive mood and behavioural changes, progressive motoric changes and

progressive neurologic changes, as well as other observable concomitants of dementia. These efforts can be traced back more than 30 years to the work of de Ajuriaguerra and associates³⁴⁻³⁶, Swiss investigators who were strongly influenced by Piaget's investigations of the stages of normal infant and childhood development. More recently, Cole and co-workers in Canada employed this approach in their Hierarchic Dementia Scale³⁷⁻³⁹. A similar approach was pursued, apparently independently, by Haycox⁴⁰ in the USA and Gottfries *et al.*⁴¹ in Sweden.

The CDR staging has a "sum of boxes" approach that employs a hierarchical, multi-axial-like procedure²¹. Based upon their seven-stage GDS, Reisberg and associates also described axial and multi-axial concomitants of progressive dementia^{42,43}. Ultimately, these latter descriptions resulted in the most detailed hierarchic staging of progressive dementia proposed to date, a 16-stage measure of progressive functional change. This latter assessment, termed Functional Assessment Staging or FAST⁴⁴, has been enumerated to be optimally concordant in dementia of the Alzheimer type, with the corresponding GDS stages, discussed above. The developers of the FAST note several advantages of this measure, including: (a) the FAST is capable of describing the entire course of dementia of the Alzheimer type, ordinally (i.e. hierarchically), in unprecedented detail; (b) the scale can assist in differentiating dementia of the Alzheimer type from other dementia processes^{22,45,46}; (c) the scale can assist in identifying premature and potentially remediable functional changes in AD patients (e.g. premature loss of ambulation due, for example, to the side effects of medication)^{22,45,46}; (d) the scale permits the retrospective as well as prospective examination of the temporal course of AD^{6,7}; and (e) the scale is the only current measure that permits the detailed staging of severe AD^{3,26,47}. A strong relationship between this FAST procedure and comprehensive cognitive assessments such as the MMSE has long been noted^{48,49}. However, because the MMSE and other cognitive modalities bottom out prior to the final five to eight FAST substages, complete concurrent validation and examination of the FAST had to await the development of cognitive measures useful in most severe dementia. Such measures were developed towards the end of the twentieth century and do, in fact, evidence strong relationships with the final FAST stages³. Subsequent work showed equally strong relationships between this latter portion of the FAST and ostensibly cognition-independent neurologic reflex changes²⁷ as well as hippocampal neuropathologic changes in volume⁶, cell number⁷, and neurofibrillary changes⁷ with the advance of AD as per the FAST stages and substages. The correlation between the advance of AD as measured with the FAST in the latter portion of the course of AD, generally after the MMSE bottom (zero) point, and cognitive, neurologic reflex and neuropathologic hippocampal measures are generally approximately 0.8-0.9, comparable to the correlation between the FAST and the MMSE in the MMSE sensitive portion of the FAST staging assessment. Because of these properties and others, the FAST, as well as the GDS staging more generally, has proved of widespread utility. For example, in the USA, the FAST staging procedure is currently utilized as the Medicare mandated "gold standard" for hospice admission (Health Care Finance Administration, 1998) and the FAST and GDS have been utilized to survey the severity mix in the US National Institute on Aging Special Care Units consortium⁵⁰. Importantly, because of the sensitivity of the FAST staging procedure to the course of AD, not only over the final seventh stage when the MMSE is zero, but also over the course of FAST stage 6, when patients still score on measures such as the MMSE, the FAST measure has shown a significant neuroprotective effect on course in a multicenter drug trial, whereas the MMSE was not sensitive to change in this study².

SUMMARY

Staging can be useful in identifying potential treatments for AD and other dementias, as well as in assessing the course and the management needs of the dementia patient, and also in the diagnosis and differential diagnosis of dementing disorders. In providing an overview of the course of dementias such as AD, from the initial to final clinical symptoms, staging is uniquely useful. Staging is also uniquely useful in assessment at various specific points in the evolution of dementing processes. Very importantly, staging can uniquely relate to management needs and the general management import of dementing processes.

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Psychogeriatric Assessment Scales

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The Psychogeriatric Assessment Scales (PAS) aim to assess the clinical changes of dementia and depression using a set of continuous scales¹. The PAS consists of two parts, an interview with the patient and an interview with an informant who knows the patient well. The purpose of interviewing both the patient and an informant is to acquire different perspectives on the patient's impairments. The PAS consists of six scales, three derived from the interview with the patient and three from the interview with the informant. The content of these scales is as follows:

Patient Interview:

- *Cognitive impairment.* This consists of brief tests of cognitive functioning and is sensitive to dementia. It correlates highly with the Mini-Mental State Examination.
- *Depression.* This asks about common symptoms of depression. It correlates highly with the Goldberg Anxiety and Depression Scales.
- *Stroke.* This scale asks about symptoms of cerebrovascular disease and is useful in differentiating Alzheimer's dementia from vascular dementia. It correlates highly with the Hachinski Ischemic Score.

Informant Interview:

- *Cognitive decline.* This scale asks questions about everyday cognitive functioning and is sensitive to dementia. It correlates highly with the Informant Questionnaire on Cognitive Decline in the Elderly (IQCODE).
- *Behaviour change.* The questions in this scale cover aspects of behaviour that could cause interpersonal difficulties and are sensitive to both depression and dementia.
- *Stroke.* The scale involves the same questions as asked of the patient, but provides an independent source of information.

Each scale yields a score along a continuum, and norms are available to show how rare a given score is in the population. The scores can be plotted as a graph to give a readily interpretable summary of the patient's pattern of impairments.

The PAS scales were derived from a principal component and latent trait analysis of items that were designed to cover the ICD-10 and DSM-III-R diagnostic criteria for dementia and depression¹. Items for the PAS scales were selected to have steep slopes (i.e. to be highly discriminating items) and to have a range of thresholds (i.e. to cover a range of severity).

Validity has been assessed against clinical diagnoses of dementia and depression using receiver operating characteristic (ROC) analysis^{1,2}. The Cognitive Impairment and Cognitive Decline scales perform well as screening tests for dementia, while the Depression scale performs well as a screening test for depression. The Behaviour Change scale is non-specific, being affected by both dementia and depression. The Stroke scales perform well at discriminating vascular from non-vascular (mainly Alzheimer's) types of dementia.

The PAS materials and User's Guide³ can be downloaded free from the Web at <http://www.mhri.edu.au/pas/>. For people who do not have Internet access, printed copies are available by writing to: PAS Project, Centre for Mental Health Research, Australian National University, Canberra 0200, Australia. The PAS is available in a number of other languages, including French, German, Italian, Chinese and Korean. For details on their availability, write to the above address.

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Computer Methods of Assessment of Cognitive Function

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Almost 20 years ago, the Royal College of Physicians recommended the use of automated testing procedures in assessing the deficits of patients with dementia, particularly in the context of clinical trials, because of their greater reliability and objectivity¹. Since then, there have been several developments that have capitalized on the explosion of computer technology and its general availability over the past decade, which have been the subject of earlier reviews²⁻⁵.

The advantages and disadvantages of computerized testing can readily be summarized. Apart from the obvious objectivity and accuracy of the measures and the standardization of the administration of the tests themselves, there are potential gains in patient compliance. Paper and pencil tests and clinical assessment interviews are generally admitted not to be popular with patients, perhaps because of their confrontational and formal nature. In our experience, computerized tests are also preferred by clinical assessors, who find that they have more time to focus on the patient during testing, rather than upon the presentation of the test material and data recording. Well-designed computerized tests that provide adequate feedback may provide some incentive for patients to do well, thereby avoiding the difficult problems of interpretation provided by lack of motivation. On the other hand, there is often little to be gained from the mere administration of standard tests in a computerized format, except in terms of the logging and storage of the data. Certainly it is infeasible to dispense with the human assessor during computerized testing; indeed, it is vital that a trained individual is present during such testing. And requesting patients to interact with certain novel and complex forms of interface, such as keyboards and other complex manipulanda, may serve only to confuse assessment by providing the patients with an extra set of problems to master. In recent developments, these problems have been circumvented, at least in part, by the use of touch-sensitive screens, which require the patient only to respond directly to the test stimuli themselves without imposing further demands by requiring them to divide their attention between a screen and a keyboard. It can also be argued that computerized assessment removes some of the creative flexibility that an experienced clinical neuropsychologist can bring to patient testing; however, this, too, can largely be avoided by the use of sufficiently flexible computerized contingencies.

In this brief review, we will concentrate on some prominent batteries that have been used for assessing cognitive functioning in the elderly. These batteries share some common principles, but also emphasize different aspects of design and administration.

EARLY ATTEMPTS: A COMPUTERIZED EVERYDAY MEMORY BATTERY

Crook and his colleagues took into account clinical, theoretical and psychometric considerations in designing their battery which, however, no longer appears to be generally available. However, their design historically represented a methodological advance in several respects and so we describe its main features below. In particular, they addressed the problem of ecological validity by simulating everyday memory situations, such as memory for faces, the locations of objects, telephone numbers and shopping lists, narrative memory for a simulated news broadcast and topographical memory for routes⁶⁻⁸. This required at that time quite advanced computer technology, including the use of the touch-sensitive screen, laser disk and video recordings. Thus, for example, one test involves viewing a live colour video recording of people introducing themselves. The subject has to learn the names that go with the faces and retain them over a 40 min delay. Tests of immediate memory for telephone numbers are administered with the subject actually dialling numbers on a telephone linked to the computer, having read a seven- (or 10-) digit number from the monitor screen. A reaction time task is configured to resemble the familiar situation of having to stop and start a car in relation to the colour of the prevailing traffic lights. The effects of certain theoretically interesting variables, such as degree of interference, can also be simulated, for example, by having the subjects hear an "engaged" signal and then asking them to redial the number.

As well as achieving a clear face validity, the battery developed by Crook and his colleagues had a degree of theoretical or construct validity. Recognizing that many of the classical memory batteries, such as the Wechsler Memory Scale⁹, in fact have a complex factorial structure including dimensions that can be labelled Attention/Concentration and Orientation, as well as Immediate and General Memory, the battery contains a more comprehensive examination of information-processing capacity that includes, for example, explicit measures of speed of reaction. In addition, evidence for dissociable forms of memory process¹⁰⁻¹¹ led to the inclusion of tests that probed different aspects of memory.

Although unfortunately this battery no longer appears to be in general use, it was potentially useful in the diagnosis of various age-related memory impairments, including those resulting from Alzheimer's disease, and in the evaluation of candidate pharmacological treatments. It was mainly used to study the relationship between ageing and memory loss in a

sample of over 3000 healthy normal subjects in the USA⁶. This study revealed some interesting differences among the tests. For example, the name-face association task administered to 1547 individuals showed a decade-by-decade decline in performance beginning in mid-life, whereas a facial recognition paradigm revealed decline only in the 70+ group in another survey of 326 subjects. Moreover, in the latter study, performance was most impaired at a delay interval of 0 seconds, suggesting attentional rather than mnemonic dysfunction.

The most important theme of the Everyday Memory Battery, which still makes it of contemporary interest, is its adherence to the principle of ecological validity—cognitive functions are tested in the context of everyday functions, which makes them relevant to patient and clinician alike. This is clearly an important element of assessment, but it should also be pointed out that it may be at the expense of test sensitivity, inasmuch that patients can resort to a greater extent on well-learned routines, to overcome deficits that would be exposed by more abstract or unfamiliar test material. In this sense the following two batteries to be considered place less weight on ecological validity and more on theoretical issues, such as the nature of the brain systems damaged in dementia, as well as on practical considerations that enhance the sensitivity and stability of the results obtained.

CANTAB (CAMBRIDGE NEUROPSYCHOLOGICAL TEST AUTOMATED BATTERY)

CANTAB was developed by a group including Dr T. W. Robbins at the University of Cambridge and Dr B. J. Sahakian (Section of Old Age Psychiatry, Institute of Psychiatry) from a research programme funded by the Wellcome Trust to improve the comparative assessment of cognition from animals to humans. CANTAB is a set of computerized neuropsychological test batteries with three main components^{4,13}:

1. *Visual memory*. Consists of short pattern and spatial recognition memory tests, a simultaneous and delayed-matching-to-sample test, and a test of paired-associate, conditional learning of pattern–location associations¹⁴.
2. *Attention*. Consists of tests of intradimensional and extradimensional set-shifting (analogous to the Wisconsin Card Sorting Test), a reaction time-based visual search task for conjunctive features, according to the Sternberg paradigms¹⁵, and a test of sustained attention, termed rapid visual information processing.
3. *Spatial working memory and planning*. Consists of tests of spatial span, spatial working memory and a computerized version of the Tower of London task, which includes separate measures of thinking and movement time¹⁶.

Each battery employs a touch-sensitive screen and begins with preliminary tests of sensorimotor function, which enable subjects with marked visual or motor deficits to be screened out from subsequent testing. The theoretical rationale for the tests is based on two major themes: first, those animal tests of cognitive function (e.g. delayed matching to sample, spatial working memory and the attentional shifting task, which is based on animal learning theory) that have proved useful in establishing the neural substrates of certain types of cognitive function and can be adapted appropriately for human subjects; and second, a componential analysis of cognitive function, allowing the characterization of elementary processes contributing to cognition. This is exemplified in each battery, e.g. visual memory, where there are separate tests of spatial and pattern recognition memory, as well as a test in which patterned information has to be remembered in specific spatial contexts. In the attentional battery, the attentional shifts are preceded by stages establishing

the subject's capacities for simple rule use, acquisition of a simple rule (simple discrimination), its reversal, and its application to more complicated stimuli with additional perceptual dimensions. In the planning battery, some of the componential cognitive requirements for the Tower of London planning task itself, such as the capacities of subjects to remember and use a short spatial sequence, and to employ working memory in a strategic spatial search task, are separately measured. In addition, a "yoked motor control" is employed, which allows the computation of thinking time, corrected for any problems of movement that inevitably confound the interpretation of latency measures in brain-damaged and elderly subjects^{16–17}. This yoked control is available to capitalize on some of the advantages conferred by computerized testing; the exact sequence of moves that a patient uses in attempting to solve a problem is stored on-line, and played back to him/her one at a time, in the yoked control test.

The "componential" approach also has some more practical benefits. For example, each test, in addition to containing internal controls, begins at a simple level, so that virtually all subjects achieve a score above floor levels. Moreover, if successful, a subject proceeds to difficult versions of the same test, which avoids ceiling effects. In addition, as the tests are non-verbal in nature, they avoid problems posed by specific language disorders and by illiteracy, and this also makes the tests suitable for cross-national studies. Finally, the tests are available for the IBM machines or similar clones running under WINDOWS '95 or later, which are relatively inexpensive as well as portable (in fact, CANTAB can be implemented on a portable computer equipped with a touchscreen) and so can be used by moderately equipped clinical neuropsychology departments of hospitals, as well as for testing in subjects' or patients' homes^{4,13}. The tests are both comprehensive, in terms of the range of cognitive functions they cover, and also sensitive to deficits in patients with dementia of the Alzheimer type (DAT), Parkinson's disease (PD), Huntington's disease and dementia of the Lewy body type^{14–20}. Some of the deficits have shown double dissociations across these groups. For example, PD patients show deficits on matching-to-sample, independently of delay, whereas patients with DAT exhibit a delay-dependent decline in performances. In addition, DAT patients early in the course of the disease can perform better than non-medicated, early-in-the course patients with Parkinson's or Huntington's diseases in tests of attentional shifting^{15,18,19}. These results are important in showing different profiles of deficits in different forms of neurodegenerative cognitive disorders, which may prove important in the early detection and diagnosis of neurodegenerative diseases and for establishing the neural substrates of the early forms of dementia. For example, it seems likely that the impairments in visual recognition memory that occur early in the course of DAT may reflect temporal lobe deficits, whereas the impairments in attention shifting in unmedicated Parkinson's disease¹⁵ and Huntington's disease¹⁹ may depend upon fronto-striatal forms of dysfunction. Other studies have confirmed that the spatial working memory and planning tests, as well as that of attentional shifting, are sensitive to frontal (but not temporal) lobe damage in humans^{16,21}.

In the assessment of dementia the results are important because the computerized tests may prove to be more sensitive to cognitive decline than many of the existing instruments, e.g. Mini-Mental State Examination (MMSE)²² and clinical stagings of dementia²³. Follow-up testing in populations of DAT and elderly depressed patients with the visual memory battery has shown that many of the tests are sensitive to the effects of progressive intellectual decline in the DAT patients and to the effects of recovery in the depressed group²⁴. Some studies have provided evidence that one of the CANTAB tests (paired

associate learning) is capable of discriminating between those patients with questionable dementia that go on to be diagnosed with DAT, and those that do not^{25,26}. There is evidence that some of the tests are sensitive to drug treatments in DAT patients²⁷, as well as to the effects of a variety of drugs in normal subjects, including those that may provide models of different aspects of dementia^{28,29}.

CANTAB has also been subject to several forms of validation-testing. There are, for example, data on test-retest reliability showing that most of the tests fall into categories described as "good" or "fair"³⁰⁻³², even though many tap "fronto-executive" functions that are notoriously unreliable in terms of measurement in this context. Most of the elements of the CANTAB battery have been administered to a large sample of normal subjects (4800) from the North East Age Research Panel. This has resulted in two major publications, which have provided a standardization of the test scores across a wide range of ages and several levels of intelligence^{33,34}. There are also burgeoning data on developmental norms³⁵. CANTAB is also validated in theoretical terms, partly from the studies of so many patient groups, including those with specific damage to different regions of the neocortex^{16,21,36}. There is also a parallel CANTAB battery for testing monkeys, which potentially provides one way of achieving a vertical integration of findings across species³⁷. Further data on the underlying neural substrates of the tests is provided via the availability of functional imaging data (mainly derived from positron emission tomography) that confirm which neural networks are activated by different tests³⁸⁻⁴².

COGDRAS: A COGNITIVE PERFORMANCE BATTERY

The Cognitive Drug Research Computerized Assessment System (COGDRAS) is rather complementary to the other two batteries described, in that it is based more on pragmatic considerations of assessing such functions as reaction time and elementary aspects of memory than on theoretical preoccupations with the ecological validity or the relationship of task performance to functional brain circuitry. COGDRAS was originally designed by Dr Keith Wesnes⁴³ to evaluate the cognitive effects of drugs in normal volunteers and patients. A further version of the battery (COGDRAS-D) was developed to examine cognitive performance in people with dementia^{43,44}. In COGDRAS-D eight cognitive tests are presented to the subject, the system originally having been installed on a BBC microcomputer⁴³. The subject faces the screen with two index fingers resting on two response buttons. All material is presented visually on the screen in large bold type. The tests comprise immediate and delayed verbal and picture recognition, a test of sustained attention similar to that used in CANTAB, simple and choice reaction time tests, and a memory scanning task. The battery deliberately sets out to avoid problem-solving tasks or the provision of negative feedback. In various extensions of the battery, tests of motor control are also incorporated. Like CANTAB, it has been employed in a wide variety of applications, especially for testing effects of drugs⁴⁵ or potential environmental toxins in normal subjects.

One validation study⁴⁴ for the dementia version of the battery has assessed 98 unselected patients from a memory clinic, who were divided into five groups on clinical assessment, including demented, depressed, "worried well", minimally cognitively impaired, and other brain disorders. The battery discriminated between some of these groups. In the key comparison of dementia and minimally impaired patients, 6/14 measures were significantly worse than for the demented group, although the level of significance obtained for these individual measures was less than

that achieved by the MMSE. It has also been used to compare patients with DAT and Huntington's disease⁴⁶. An earlier study⁴³ showed good test-retest reliability coefficients for demented patients, particularly on the reaction time measures and significant correlations with other commonly-used instruments in dementia research. More recent applications of the battery have also shown that it is possible to test DAT patients over as long as a 4h period, in the context of studying the effects of the benzodiazepine antagonist flumazenil⁴⁷.

CONCLUSIONS

The batteries reviewed above, as well as others, will ultimately have to be compared with one another, as well as with more conventional assessment procedures, and this will require large and dedicated studies, some of which are under way. A clinician anxious merely to diagnose DAT and other conditions will understandably enquire why such complex tests should supplant the use of such simple and easy-to-use instruments as the Mini-Mental State Examination²². The answer is that they are not intended to supplant, but rather to augment, their use. In our own experience, for example, the Mini-Mental State Examination often fails to detect the early onset of dementia, especially in individuals of high IQ. To understand the relationship between such clinical rating scales and the specific computerized tests is to consider the scale as providing a gross measure on a relatively undifferentiated range of functions, whereas the computerized tests provide more specific and precise information about particular capacities, thus providing important information for patient management.

Crook *et al.*⁶ pointed out that they found it useful to develop in parallel their own self-rating scales for memory, although the relationships with direct performance measures are typically low. Of course, such low correlations may reflect the measurement of subtly different components of memory, all of which should be taken into account. They view self-rating data as important in assessing age-related memory disorders and family ratings and clinical scales as useful in the assessment of dementia. Certainly, it is important to relate what may be a statistically significant and consistent effect of a treatment on computerized tests of cognitive performance to the clinical improvement manifest to the patient's family in their everyday activities. However, it should be noted that such assessments of everyday activities have not so far proved to be especially discriminating for patients with mild DAT. It is particularly important to add some assessments of everyday activities to the assessments provided by CANTAB and the COGDRAS battery, as these are not specifically designed to simulate such situations. The rationale is that the use of rather visual abstract material tends to be more sensitive in detecting deficits than the use of concrete examples, as there is every indication that many well-established skills (e.g. reading) or types of knowledge (skills) may remain largely intact in early dementia. Moreover, obvious differences in life experience, e.g. produced by different occupations, will tend to be minimized. Finally, we should point out that it is important that batteries such as CANTAB, and to a lesser extent CDR, can provide means for differentiating cognitive disorders in the elderly arising from a variety of conditions, preferably on a qualitative basis. Providing a profile of which functions are spared as well as which are impaired may have implications for strategies based on rehabilitation, as well as pharmacological treatments, which will increasingly motivate attempts at cognitive remediation for the elderly.

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The Assessment of Depressive States

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The diagnosis of depressive states in the elderly requires a multi-dimensional approach and should not be undertaken without a clear understanding of the normal aging process and the psychosocial factors that are unique to the elderly. The accurate assessment of depressive disorders in the elderly is essential, since this population is often subject to more adverse side effects from pharmacotherapy, and concurrent depression can limit compliance with medical treatments and worsen the cognitive decline associated with dementia. Further, depression can adversely affect the clinical course of other medical illnesses and increase morbidity and mortality^{1,2}. Depression in late life is not a normal part of aging and is not necessarily more difficult to treat or chronic when compared to depression in younger people^{3,4}.

A number of authors have drawn attention to the psychosocial stressors that must be taken into account in the diagnosis of depression in the older population and to the high levels of depressive symptoms in the elderly living in the community⁵. Although the prevalence of individuals with major depression may not be as high as for younger subjects in such settings^{6,7}, the prevalence increases in elderly patients in long-term care facilities and the medically ill^{8,9}. Depressive symptoms can lead to accelerated cognitive decline, physical impairment and increased health care costs. The elderly population maintains the highest suicide rate^{10,11} so that the social and economic consequences of this disorder are severe.

THE SYNDROME OF LATE-LIFE DEPRESSION

To accurately assess the depressed older adult, the clinician must be aware of the major syndrome encountered in older adults. The major categories of mood disorders outlined in the *Diagnostic and Statistical Manual of Mental Disorders*, 4th edn¹², are major depression, dysthymic disorder (dysthymia) and adjustment disorder (bipolar disorder is discussed elsewhere), which correspond roughly with the International Classification of Diseases 9 (ICD-9)¹³ for diagnoses of manic-depressive psychosis, neurotic depression and brief depressive reaction, respectively. The revision of the ICD (ICD-10)¹⁴ has diagnoses of depressive episode and persistent affective state (with a subclassification of dysthymia) that closely approximate the corresponding DSM-IV classifications.

Major depression is distinguished by the severity of symptoms, which may include significant weight loss or weight gain, insomnia or hypersomnia, psychomotor agitation or retardation, diminished ability to think or concentrate, feelings of worthlessness or excessive inappropriate guilt and recurrent thoughts of death or

suicide¹². A melancholic depression (DSM-IV) or severe depression (ICD-10) represents a subtype of major depression that is felt to be preferentially responsive to somatic therapy (i.e. antidepressants and electroconvulsive therapy) and includes symptoms such as a lack of reactivity to pleasurable stimuli, diurnal variation in mood with depression worse in the morning, early morning awakening, psychomotor retardation or agitation, significant weight loss and other factors that may have a bearing on response, including no significant personality disturbance before the onset of a major depressive episode and a previous history of a major depressive episode that may have responded to somatic treatment. In the DSM-IV, depression is sub-typed according to whether the mood disturbance is associated with a seasonal variation (e.g. a temporal relationship between a distinct 60-day period of the year) or psychotic symptoms that are more likely to be mood-congruent (i.e. delusions or hallucinations of a depressive nature). Psychotic depressions may be more common in late life¹⁵.

The DSM-IV now includes a research diagnosis of minor depression. The duration of the episode and the symptoms are the same as major depression. However, in minor depression the number of symptoms required for diagnosis is less compared to major depression. Minor depression is important in the elderly as there is evidence that these individuals have levels of cognitive and functional impairment similar to those who met full criteria for major depression¹⁶.

Dysthymic disorder (dysthymia) is defined as a chronic low-grade depression, which lasts at least 2 years (DSM-IV), or "several" years (ICD-10). Primary dysthymia may occur in older adults due to changing roles and life conditions. Personality styles and the individual's ability to cope with changing life situations may predispose the individual to develop a dysthymic disorder (see Chapters 71, 74). Although certain severe personality disorders are less common in the elderly, long-standing patterns of perfectionism and the need for external gratification can lead to chronic low self-esteem and dysphoria. Dysthymic disorder may also develop secondary to a medical disorder, which leads to chronic debilitation or other psychiatric disorders, such as substance abuse, anxiety disorder or somatization disorder.

The elderly patient may also have to adjust to severe changes in lifestyle and the loss of loved ones. In the DSM-IV, the patient is not classified with an adjustment disorder unless his/her reaction is considered maladaptive and the level of impairment is significantly severe, so as to be greater than what would be normally expected. The ICD-10 outlines similar symptoms under the heading of mild depressive episodes. Several common life events that prove to be stressors in the elderly

include retirement, financial problems, crime rates, physical illness, the death of a spouse and moving to an institutional setting. Whether the elderly subject's reaction to these stressors is classified as a disorder is a clinical judgement. Regardless, the clinician should watch carefully for the development of symptoms of a major depression if the individual's reaction to stress is persistent or severe.

The symptom profile of older patients with depression has also been posited to differ from that of younger adults, and these subjects are more likely to demonstrate hypochondriacal or somatic symptoms. However, Blazer asserts that this issue is confounded by the significant co-morbidity with other physical illnesses, and these factors must be controlled when making such comparisons³. Hypochondriasis, or the belief that one has a serious physical illness, is quite different from an older patient's complaints of physical symptoms such as constipation and nausea. In the hypochondriacal patient these complaints are more likely to be long-standing and to fluctuate over time. In the depressed patient, somatic complaints may become severe and delusional. Somatic delusions can be bizarre and unshakeable. Whereas the patient with hypochondriasis is willing to consider a physical cause for the symptoms, the depressed patient is more likely to have a bizarre rationalization for his symptoms that defy logical explanation. The depressed patient is also apt to show a history of neurovegetative symptoms that have become progressively worse over time.

THE PROBLEM OF CO-MORBIDITY

The diagnosis and treatment of major depression in older adults is more often complicated by concomitant medical illness and/or cognitive decline than in their younger counterparts. For example, over 10% of patients who are initially diagnosed with dementia were found to have an affective disorder when seen in follow-up¹⁷⁻¹⁹. This misdiagnosis is due to the fact that the symptoms seen in dementia, such as withdrawal, diminished ability to think and concentrate and psychomotor retardation, occur in both dementia and depression. In fact, over 50% of patients with dementing illnesses such as Alzheimer's disease may also have depressive symptoms, with 20% meeting criteria for a major depressive episode²⁰.

Alexopoulos *et al.*²¹ demonstrated that a significant minority of individuals with depression and reversible cognitive deficits eventually progress to true dementia within 3 years. Krishnan *et al.*²⁴ found an association between apolipoprotein E-ε3/ε4 (which has been linked to Alzheimer's disease^{22,23}) and major depression in later life, further supporting the association of dementia and late-onset depression²⁴.

Various clinical studies have shown that depressed older subjects are less likely to demonstrate depressed mood²⁵ or guilt²⁶ and more prone to complain of fatigue²⁷. Other researchers emphasize that these differing presentations are the result of accompanying dementia and medical illness that are more likely to occur in older depressed patients, and that depressed elderly patients without concomitant medical illnesses have presentations similar to younger subjects²⁸.

Support for this latter view has been provided by Blazer and his colleagues²⁹, who compared depressive symptoms in a group of middle-aged (35-50 year-old) and older (>60 years) inpatients with melancholic depression. The symptom profiles of these patients were markedly similar to the elderly, differing only in their more frequent reporting of weight loss and less frequent suicidal thoughts. Finally, Blazer discusses some of the misconceptions of depression in the elderly, pointing out that as discussed above, symptomatically there is little difference when compared to adult early-onset depression.

TAKING A HISTORY FROM THE DEPRESSED OLDER ADULT AND FAMILY

The key elements in the patient's history include evidence of a previous psychiatric illness especially a major depressive or manic episode, particularly if that episode responded to somatic therapy (e.g. lithium, antidepressants or electroconvulsive therapies). Obtaining pharmacy and other medical records regarding the symptoms, dose and the duration of medication trials can further aid in the assessment of depressed individuals. Major depression is a recurrent illness, with about half of the patients experiencing two or more episodes in their lifetime. In order to distinguish dementia from depression, researchers have relied upon the findings in the family and individual histories, mental status and laboratory examinations. Determining the time of onset of the depressive symptoms may also be useful, since most dementias are slowly developing, with early evidence of cognitive decline (e.g. the inability to balance a checkbook) being present months to years before the typical form of the disease becomes apparent. In contrast, the course of depressive illness may seem more abrupt.

An accurate family history may also provide valuable diagnostic information, since dementing illnesses (e.g. Huntington's chorea and Alzheimer's disease, particularly the early-onset form) and major depression (especially the early-onset form) have both been demonstrated to have a genetic diathesis. The family history should also include questions about neurological and medical disease, alcohol/drug abuse, anxiety, suicide and psychosis, since these conditions are often associated with concomitant depression.

THE MENTAL STATUS EXAMINATION

The mental status examination and standardized depression rating scales may also aid in determining whether the patient has depression, dementia or (as is often the case) both.

The depressed patient approaches the mental status examination with the same lack of interest that is also apparent in other areas of his/her life. The depressed patient may attempt to prematurely terminate the interview or consistently refuse to try and answer questions. On the other hand, the demented patient may be quite concerned about his/her declining cognitive skills and, at the same time, make attempts to cover their deficits with confabulation³⁰. He/she may use notes and other reminders to keep up with facts and uncovering his/her deficits might be quite distressing. This is in marked contrast to the apathetic attitude of the depressed patient. The depressed inpatient is also more likely to be able to find his/her way around the ward and to keep track of the ward routine and personnel. In contrast, the demented patient will forget mealtimes and confuse familiar faces. This confusion may often get worse as evening approaches and is incorporated in the well-known clinical syndrome of "sundowning".

The mental status examination of patients with physical illnesses may show primarily problems with concentration and attention. In more severe cases, evidence of delirium may be present. Delirium is diagnosed when the level of consciousness becomes impaired. In this condition, disorientation and memory impairment may be present along with illusions (which are misinterpretations of environmental stimuli), hallucinations and disturbances in psychomotor activity and the sleep-wake cycle. As opposed to dementia, these symptoms usually evolve over hours and days rather than months and may show a waxing and waning course along with autonomic instability.

A common clinical dilemma occurs in patients who are known to be demented, but in whom the clinician seeks to make an accurate diagnosis of depression. In these patients, the clinician may need to assign more weight to neurovegetative signs, such as weight loss

and insomnia, although these same symptoms may occur in non-depressed demented patients if they are too disorganized to prepare a meal or have night-time confusion. Some researchers have pointed to the presence of cognitive symptoms of depression (i.e. depressed mood, anxiety, helplessness, hopelessness and worthlessness) as being more prominent in demented patients, whereas neurovegetative signs are notably absent³¹.

Monitoring the patient over time, observing for some overall consistent pattern of depressed mood and affect, is also helpful. Although there is little or no data to support empiric trials of antidepressants, given the relatively low side-effect profile of the newer antidepressants³² (SSRIs: see Chapter 78), many clinicians will often attempt a trial of an antidepressant in these difficult situations in hopes the treatment may improve cognition and function.

PSYCHOLOGICAL TESTS

Standardized assessments of cognitive performance (e.g. Wechsler Adult Intelligence Scale and Clinical Dementia Rating Scale³³) may also be useful in distinguishing depression and dementia. The depressed patient will more likely show results that are inconsistent over time and are effort-dependent. The demented patient will demonstrate a more global decline, with relatively lower scores on aspects of the testing requiring adaptability and processing of information (e.g. performance IQ) as opposed to rote skills and long-term memory tasks (e.g. verbal IQ).

Standardized depression scales may yield false-positive results in demented patients, particularly if they are heavily weighted for difficulties symptoms, such as with attention and concentration, which overlap with symptoms of depression. Depression scales that have been specifically designed to grade levels of depression in patients known to be demented include the Cornell Depression Scale³⁴ and the Dementia Mood Assessment Scale³⁵. The Geriatric Depression Scale (GDS), Beck Depression Inventory (BDI) and the Montgomery–Asberg Depression Rating Scale (MADRS) are other useful tests that can help aid in evaluation of a depressed patient^{36–38}.

LABORATORY TESTS

The common laboratory tests for depression may lose much of their specificity in demented patients. The Dexamet has one suppression test that is frequently abnormal in dementia^{39,40}.

The sleep electroencephalogram, however, does maintain its specificity and is able to distinguish a group of elderly depressed patients from those with dementia^{41,42}.

Any physical illness that can potentially affect the central nervous system can present with depressive and dementing symptoms; these illnesses include encephalitis, chronic subdural hematoma and normal pressure hydrocephalus. Depressive symptoms such as fatigue, insomnia, weight loss and concentration problems can occur in other illnesses that do not directly affect the central nervous system (CNS). These illnesses include infectious diseases (e.g. tuberculosis, influenza), cardiovascular disease (e.g. congestive heart failure), endocrine abnormalities (e.g. thyroid disease, diabetes), electrolyte disturbances (e.g. hyponatremia, hypocalcemia), renal and hepatic disease.

Recent evidence has indicated that both cerebral vascular accidents and periventricular⁴² white matter disease are related to depressive symptoms in the elderly^{44,45}.

MEDICAL WORK-UP

Medical conditions also present with depressive symptoms in the elderly. Many of these medical conditions are reversible with

Table 29.1 Laboratory work-up in the depressed elderly patient

Laboratory test	Underlying condition
Complete blood count	Infectious disease Encephalitis Meningitis Sub-acute bacterial endocarditis Anemia
Electrolytes	Hypokalemia Hyponatremia Diabetes Uremia
B ₁₂ /Folate	Pernicious anemia
Thyroid panel	Hypo/hyperthyroidism
Liver enzymes	Hepatitis Liver cancer
Calcium/phosphorus	Hyperparathyroidism
Guaiac stool	Colon disease
Urinalysis	Infection Renal disease
Consider additional test including:	Particularly patients who complain of fatigue, edema, shortness of breath or who are being considered for antidepressant therapy
Electrocardiogram	
Electroencephalogram	When delirium may be a concern or to screen for a mass lesion
Urine drug screen	For lead (exposure to paint), mercury (textile manufacturing), organophosphates and arsenic (insecticides). Also drugs of abuse (e.g. opiates, marijuana, etc.)
Human immune deficiency virus (HIV)	Exposure to potentially contaminated blood products (surgery, drug use) or a history of promiscuity or homosexuality
Brain computed tomography	Neoplasia, stroke
Brain magnetic resonance imaging	Particularly to view the posterior brain stem

proper treatment, and estimates of the number of patients with reversible conditions presenting as dementia have been as high as 15%¹⁷. The work-up of medical conditions that present with depressive symptoms in the elderly includes a personal history of medical illness, mental status examination and laboratory tests⁴⁶.

The physical examination and laboratory diagnosis is crucial in the diagnosis of medical disorders. The typical laboratory screening for elderly patients with depressive symptoms is outlined in Table 29.1. Clearly, the medical history and physical examination would influence the laboratory testing that was actually done. The neurologic examination should be particularly thorough and include an examination of both subtle (e.g. frontal reflex signs) and prominent (e.g. reflex changes) signs of neurologic dysfunction.

MEDICATION HISTORY

Medication, including the antihypertensive medications such as propranolol and centrally acting α -methyl dopa and reserpine, has been shown to cause depressive symptoms. Drugs such as alcohol are also frequent causes of depression in the elderly, as are the barbiturates that are used as sedatives. Patients at risk should be screened for heavy metals, including lead, arsenic and the organophosphates.

SUMMARY

Depressive symptoms are common in the elderly, whereas the syndrome of a major depressive episode is relatively rare^{6,45}.

Symptoms of depression can be manifestations of either dementing or medical illnesses, both of which are common in the elderly. Psychosocial factors that are unique to the elderly may be etiologic in the genesis of depressive symptoms, but nevertheless require the clinician to assess each patient fully and treat as appropriate. Accurate treatment mandates as detailed history, often with collaboration with the family, and a thorough medical evaluation for other co-morbid illnesses that may contribute to the patient's current presentation. The treatment of depressive symptoms in the elderly requires an understanding of both the origin of these symptoms and the proper treatment of the underlying disorder, whether it be due to medical, dementing, psychosocial or melancholic factors.

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The Geriatric Depression Scale: Its Development and Recent Application

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Developed in 1983, the Geriatric Depression Scale (GDS) is a 30-item self-report measure for rating depression in elderly adults¹. The identification of depression in older adults is of particular importance to clinicians, since depression is the most common psychiatric disorder in this population and can be associated with morbidity and mortality².

The GDS was designed to address the unique characteristics of geriatric depression and the subsequent difficulties in rating depression in older adults using scales developed for younger populations. Somatic complaints are less useful indicators of depression in the elderly than in young adults, since such symptoms often accompany the normal aging process. Depression in older adults is more often accompanied by subjective complaints regarding decline in memory and cognition than is depression in younger adults. Questions pertaining to suicidal intent or one's hopefulness regarding the future can be interpreted differently by elderly adults, who are in the latter stages of their lifespans. To address these issues, the GDS excludes questions pertaining to somatic complaints, registers a cognitive dimension of depression and focuses dominantly on the worries of the individual and how that person interprets his/her quality of life. Additionally, the GDS employs a yes–no format for both economy of time and ease of self-administration in this population.

The scale was developed and validated in two phases. First, 100 widely varied yes–no questions were selected and tested for their potential to distinguish depressed elderly adults from normal controls. The 30 questions correlating most highly with depression were chosen for inclusion in the final version of the GDS. Second, validity of the scale was established by comparing the mean GDS scores for subjects classified as normal, mildly depressed or severely depressed, using the Research Diagnostic Criteria (RDC) for depression. The mean GDS scores for these three groups were reliably different and were ordered in accordance with the differing RDC scores¹. The scale has been found to have high internal consistency and high test–retest reliability. A score of 11 or higher has been found to indicate the presence of depression, yielding 84% sensitivity and 95% specificity³.

The GDS has been utilized widely in clinical practice and research and has been shown to be a reliable and valid measure of depression in outpatient, nursing home and hospital settings^{4–6}.

Additionally, the GDS is sensitive to depression among elderly adults suffering from mild to moderate dementia and elderly adults with physical illnesses⁷.

To date, the GDS has been translated into 21 languages, including Spanish and Chinese. It is in the public domain, and available on the Web (<http://www.stanford.edu/~yesavage/GDS.html>). A 15-item, short form of the GDS has been developed and validated⁷. Additionally, administrations of the GDS by telephone and by requiring caregivers to answer the questions have been validated^{8,9}.

Overall, the GDS provides a sensitive measure of depression in older adults that is time-efficient, easy to administer and reliable and valid in a broad range of clinical and research settings.

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Center for Epidemiologic Studies Depression Scale: Use among Older Adults

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The Center for Epidemiologic Studies Depression Scale (CES-D) is a depression screening instrument which has been applied widely in epidemiologic studies including many community-based studies of older adults¹⁻³. The scale consists of 20 items which on factor analysis among older adults fall into four different factors. The first factor is depressed affect, and items that load on this factor include: "bothered by things that usually don't bother me", "I could not shake the blues", "I felt depressed", "I felt lonely", "I had crying spells" and "I felt sad". The second factor is positive affect and includes the items: "I felt as good as other people", "I felt hopeful about the future", "I thought my life had been a failure", "I was happy" and "I enjoyed life". The third factor is somatic complaints and includes the items: "I did not feel like eating", "I had trouble keeping my mind on what I was doing", "My sleep was restless", "I felt like everything was an effort", "I talked less than usual" and "I could not get going". The final factor is interpersonal relations and includes the items: "People were unfriendly" and "I felt that people disliked me". Each item is rated on a 1-4 scale, depending on the frequency of symptoms the week prior to the administration of the scale. Therefore, the range of answers is 0-60. A score of 16 or greater is considered indicative of clinically significant depression¹.

Murrell *et al.*⁴ found, when using the CES-D in a community of over 2000 community-dwelling adults, 55 years of age and older in Kentucky, that the mean score was 8.9 for African-Americans and 9.2 for Whites. When they applied the cut-off noted above for clinically significant depressive symptoms, 12.8% of African-Americans and 13.7% of Whites had clinically significant depression⁴. Berkman *et al.*⁵ found that 16% of both African-Americans and Whites in an urban community scored above the threshold for clinically significant symptoms. When control variables are taken into account, such as gender, socioeconomic status and functional health, the association of age and depression disappears (whereas in uncontrolled analysis there is a positive association between age and depression) using the CES-D⁶.

When using the CES-D in older adults, there is little difference by gender and little difference by race/ethnicity². In a cross-

sectional analysis, age is negatively associated with somatic complaints, lower education is associated with more complaints of depressed affect and interpersonal problems, African-American race is associated with increased interpersonal complaints and cognitive impairment associated with somatic complaints and interpersonal complaints. Of all control factors, however, disability is most closely associated with all four of the factors noted above.

In summary, the CES-D is a useful screening scale for depressive symptoms in community (and clinical) samples of older adults. The one drawback to this scale is that it requires responding along a spectrum of four responses (little or none, some of the time, most of the time, or all the time) over the previous week compared to a simple yes-no format, as is found in other symptom screening scales. The scale has been used extensively in older adults and therein lies its greatest value for future epidemiologic studies as well as clinical screening efforts.

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The Development of the EURO-D Scale

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BACKGROUND

The 11 country EURODEP consortium, assembled an unprecedented body of data; 14 population-based surveys of mental health in late life, including 21 724 older Europeans aged 65 years and over. Most centres had used either GMS/AGECAT¹ or the very similar SHORT-CARE² as their index of clinical case-level depression. However, three centres had used the Centre for Epidemiological Studies Depression scale (CES-D)³, one the Zung Self-rating Depression Scale (ZSDS)⁴ and one the Comprehensive Psychopathological Rating Scale (CPRS)⁵. Our challenge, therefore, was to derive from these instruments a common depression symptom scale, allowing risk factor profiles to be compared between centres⁶.

METHOD⁶

The instruments were scrutinized for common items. Algorithms for fitting items from other instruments to GMS were derived by either: (a) empirical observation of the nature of the relationship between items in different scales where they had been administered together; or (b) expert opinion. The resulting 12-item scale (see Table 30.1) was checked in each centre for internal consistency, criterion validity and uniformity of factor analytic profile.

RESULTS^{6,7}

The EURO-D, however derived, is an internally consistent scale. Cronbach alphas for the 14 centres ranged from 0.58 to 0.80. It seemed also to capture the essence of its parent instrument. Correlations with the CES-D in all four centres using that measure exceeded 0.90, and the correlation with the Zung was 0.84. A cut-point of 3/4 on the EURO-D scale predicted a GMS/AGECAT computerized diagnosis of depression with 70–80% sensitivity at 80–95% specificity. Principal components factor analysis demonstrated that a very similar two-factor solution seemed appropriate in all centres, however the scale had been derived; depression, tearfulness and wishing to die loaded on the first factor (affective suffering) and loss of interest, poor concentration and lack of enjoyment on the second (motivation).

In general, EURO-D scores increased with age, women scored higher than men, and widowed and separated subjects higher than others. The gender effect was negligible among the never-married but was not modified by age. In most centres EURO-D could be reduced into two well-characterized factors; affective suffering,

responsible for the gender difference, and motivation, accounting for the positive association with age.

CONCLUSIONS

Large between-centre differences in depression symptoms as assessed by EURO-D were explained neither by demography nor by the depression measure used in the survey. Consistent, small effects of age, gender and marital status were observed across Europe. Depression may be overdiagnosed in older persons

Table 30.1 EURO-D Scale.

The Geriatric Mental State (from which EURO-D is derived) is a semi-structured clinical interview. The following instructions apply:

1. Each question should be asked as it is written. The sections in parentheses are additional prompts to clarify the question if it has not been understood.
2. Sections in *italic* script provide the criteria by which the interviewer judges from the response whether the symptom is present or absent.

EURO-D item	Corresponding GMS question
1. Depression	Have you been sad (depressed, miserable in low spirits, blue) recently?
2. Pessimism	How do you see your future? <i>Pessimistic, empty expectations or bleak future</i>
3. Wishing death	Have you ever felt that you would rather be dead? <i>Has ever felt suicidal or wished to be dead</i>
4. Guilt	Do you tend to blame yourself or feel guilty about anything? <i>Obvious guilt or self blame</i>
5. Sleep	Have you had trouble sleeping recently? <i>Trouble with sleep or recent change in pattern</i>
6. Interest	What is your interest in things? <i>Less interest than is usual</i>
7. Irritability	Have you been irritable recently?
8. Appetite	What has your appetite been like? <i>Diminution in the desire for food</i>
9. Fatigue	Have you had too little energy (to do the things you want to do)? <i>Listlessness or subjective energy restriction</i>
10. Concentration	How is your concentration? <i>Difficulty in concentrating on entertainment or reading</i>
11. Enjoyment	What have you enjoyed doing recently? <i>Almost nothing enjoyed</i>
12. Tearfulness	Have you cried at all?

because of an increase in lack of motivation that may be affectively neutral, and is possibly related to cognitive decline.

The EURO-D scale shows promise as a means of harmonizing data from studies and perhaps trials that have used similar depression outcome measures. In the EURODEP datasets the EURO-D has been used informatively in three ways⁷:

1. To compare EURO-D item prevalence between centres.
2. To compare EURO-D scale distribution between centres.
3. To compare effect sizes for associations between risk factors and EURO-D score between centres.

Researchers might consider using the EURO-D in epidemiological or health services research, either on its own or as a component part of the GMS. Its principal advantage at present is the large amount of normative data available from the many population-based studies in different parts of the world to have used the GMS. A limitation of the scale is its current format, which relies upon interviewer administration and rating. A future priority will be to develop self-report forms and test their equivalence against the current version.

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Interviews Aimed at Differential Psychiatric Diagnosis

GMS–HAS–AGECAT Package

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For epidemiological and other studies of mental illness and morbidity in older age it is important to ensure as far as possible that the differences in the levels of cases of illness found between geographical areas and between studies at different points in time are not due to methodological differences and, in particular, the way the diagnoses themselves are made. To overcome this problem, standardized interviews were introduced. The GMS–HAS–AGECAT Package consists of a series of interviews designed to be given to a subject and his/her informant for assessing the dementias and depression, with optional sections for minor mental illness^{1,2}. The Geriatric Mental State (GMS) was derived originally from the Present State Examination³ and the Mental Status Schedule⁴. Substantial modifications and additions were incorporated to make it more applicable to older populations and to increase emphasis on organic states. The Package now provides, in addition to the GMS, which is an interview with the subject, the History and Aetiology Schedule (HAS) for an informant, which allows the assessment of onset and course of illness, past history, family history and certain risk factors for dementia, depression and other mental illness. The Secondary Dementia Schedule provides a semi-structured framework to aid in collecting information required for the NINCDS–ADRDA (National Institute of Neurology and Communicative Disorders and Stroke–Alzheimer’s Disease and Related Disorders Association) criteria⁵ and the assessment of daily living.

Standardization of diagnosis is achieved by the AGECAT (Automated Geriatric Examination for Computer Assisted Taxonomy) computer-assisted differential diagnosis^{6–8}. Based on an extensive decision tree method, the system aggregates the data into scores and allots each subject to levels of diagnostic confidence on each of eight diagnostic syndrome clusters, organic, schizophrenia/paranoid, mania, depression (psychotic and neurotic type) (levels 0–5), obsessional, hypochondriacal, phobic and anxiety neurosis (levels 0–4). Levels 3 and above are what would usually be recognized by psychiatrists as cases of illness. The computer then compares these levels with one another to derive a final differential diagnosis and flags cases where the decision has been difficult. The validity of the AGECAT diagnosis has been assessed against psychiatrists’ diagnoses on the same patients. The range of kappa values for the agreement between AGECAT and psychiatrists’ diagnoses for organic states is 0.80–0.88 and for depressive states 0.76–0.80⁶. Outcome studies are now providing additional validation. After 3 years’ follow-up, over 83% of

AGECAT cases of organic disorder identified in a community study were either dead or still dementing. One-third of depressed cases were also depressed 3 years later⁹. Post mortem validation studies are in progress.

In the second stage AGECAT uses the data from the HAS to take the diagnosis to a further stage, dividing organic states into acute or chronic, and the latter into the different types of dementia using a standardized form of the Hachinski score¹⁰. It also identifies bereavements and flags coexistent immobility, pain, life-long intellectual function and physical illness.

The GMS can be used by trained lay workers and provides a diagnosis by AGECAT. When used in epidemiological studies it is possible to derive prevalence figures for the full range of psychiatric morbidity using a one-stage design¹¹. The measures are therefore economical as well as reliable and valid. The Package does not rely on special psychological tests as these are not applicable across cultures or socioeconomic groups or with populations of varying literacy. The interviews have also been transferred for presentation on laptop computer, which improves accuracy and communication, avoids delays and costs in inputting data, and provides rapid access to results and easy quality control of interviewing techniques¹².

These measures have been used in a number of projects, including the Medical Research Council (UK) ALPHA^{13,14} study of the incidence of the dementias and the multicentre Cognitive Function and Ageing Study, the EURODEP EC-funded Concerted Action¹⁶, the ongoing ASIADEP studies in 12 Asian centres and the 10/66 Club studies on the prevalence of dementia in India, Latin America and Africa, as well as forming part of the minimum data set required by the EURODEM (EC Concerted Action on Epidemiology and Prevention of Dementia)^{17,18}. The GMS has been translated and used in a wide range of languages. Recently, algorithms have been developed for ICD-10, DSM-III-R and DSM-IV and compared favourably with psychiatrists’ diagnoses using these international criteria. The HAS has been shortened and modified to provide for the criteria of Lewy body dementia and other recognized dementia classifications.

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CAMDEX

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The Cambridge Examination for Mental Disorders of the Elderly (CAMDEX) was devised to assist clinicians and epidemiologists to diagnose dementia, and mild dementia in particular, as reliably and validly as possible¹. It was developed with the following aims in mind: it addresses all elements of current diagnostic criteria; it incorporates historical material so that persons with cognitive deficits due to intellectual disability, sensory handicap or functional mental disorder are classified correctly; and its neuropsychological battery is sensitive to mild dementia.

The schedule is fully structured and incorporates: a mental status examination; a comprehensive neuropsychological battery (CAMCOG); a medical and psychiatric history; a brief physical examination; and an interview with an informant that enquires into changes in memory, intellect, personality, behaviour and self-care. Medications, laboratory investigations and imaging are all recorded. There is a clear focus on dementia but conditions such as delirium, anxiety, depression, bipolar disorder, delusional disorder and schizophrenia are considered as differential diagnoses. The respondent interview takes 30–60 minutes to complete. Informants are questioned for another 20–30 minutes, by telephone if necessary. Diagnoses are based on all available data using criteria virtually identical to those in ICD-10. Since judgement is required, interviewers should have a clinical background and have received training in formulating complex data, applying diagnostic criteria and rating dementia severity.

CAMDEX, which is available in English, Dutch, French, German, Italian, Spanish and Swedish versions, is practicable and

acceptable and can be administered with high inter-observer reliability¹. It includes the Mini-Mental State Examination², Hachinski Ischaemia Scale³ and other commonly-used rating scales. When applied in community surveys, prevalence rates of dementia based on CAMDEX assessments are close to those reported in other recent Western European studies⁴. CAMDEX diagnoses are also stable over time. In a British study in which community residents rated as having mild dementia were followed for 2 years, diagnoses were maintained for 51 of the surviving 56 persons. Two were re-classified as having minimal dementia and only three were rated as having no significant impairment, mostly as the result of better-controlled diabetes mellitus^{5,6}. CAMCOG has been used as a stand-alone assessment package in many clinical and community studies. Further details about its psychometric properties can be found on the website listed below.

An updated version of CAMDEX includes both DSM-IV and ICD-10 diagnostic criteria and provides better coverage of more recently described conditions, such as frontotemporal dementia and dementia of the Lewy body type. A floppy disk permits computer-assisted administration, data entry, scoring and analysis. CAMDEX is copyright but packs including the interview schedule, test materials and scoring sheets can be purchased in most countries. Consult this website (<http://www.iph.cam.ac.uk/camdex-r>) for further updates, access to a dedicated bulletin board, references to published reports based on CAMDEX and CAMCOG, and other relevant information.

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CAMDEX

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Assessment of Daily Living

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The need for assessing activities of daily living¹ (ADL) has increased during the last few decades for several reasons. First, the number of disabled persons in the population is higher today than ever before, since more people survive longer and have to live with the consequences of diseases and other² disorders. Second, the costs for ADL-related services to disabled persons have surged ahead, both for sheltered living and for assistance given by people paid to provide services in the homes. Relatives and neighbours will often give voluntary help but they, too, may need professional support. Third, there is a need for outcome measures, other than mortality and morbidity, which can describe levels of function among elderly people. This is required for the evaluation of pharmacological treatments and rehabilitation programmes. Many people in different professions are involved in the care of the elderly, e.g. physicians, nurses, occupational therapists, physiotherapists and social workers. In clinical practice, they need a common language to communicate their knowledge about the individual's functional level. Although much work has already been done to develop new ADL instruments and make existing ADL assessments representative, reliable and valid, much remains to be done to reach a consensus in this field. Even a definition of the concept of ADL can still be a matter for discussion: "ADL" means "activities which are common to all human beings and which must be performed regularly in order to live an independent life". This is a very broad definition, which permits a variety of activities to be included. These activities are usually divided into several categories:

1. *Personal ADL*: activities concerning self-care, care of one's own body, e.g. feeding, dressing, bathing.
2. *Instrumental ADL*: activities concerning home management, e.g. cooking, cleaning, shopping.
(*Communication by talking and writing may be referred to as personal ADL, while transportation and managing money may be referred to as instrumental ADL.*)
3. *Professional work*.
4. *Leisure activities*.

Ambulation or walking is not an ADL item, although it is often assessed in this context. It is rather a level of mobility on a scale from bed-ridden, through unsteady gait such as creeping, to walking and running. A certain level of mobility is usually required for independent living as is a certain level of intellectual capacity, motivation and a suitable environment.

"ADL-capacity" means that a subject can perform ADL independently of another person. Even if a person can do things, it may not be certain that he/she actually does them in practice! In the WHO Classification of Impairments, Disabilities

and Handicaps, ADL-dependence is classified as a disability. "Disability" has been defined as "any restriction or lack (resulting from impairment) of ability to perform an activity in the manner or within the range considered normal for the human being"².

"ADL-ability" means that a subject does perform ADL. It may take time, it may hurt, or there may be other difficulties, but the person is nevertheless able to perform ADL and independently of another person. If a person does not perform ADL, it may be due to physical or intellectual capacity deficits (he/she can not or does not understand) but it may also be due to lack of motivation (he/she does not want to do it), or to environment problems (he/she is not allowed to do it or is prevented).

When assessing ADL-performance in clinical practice, it is recommended that the interviewer starts with the question: "Does the patient perform the activity or not?" If he/she does not or has difficulties, the interviewer may proceed by asking "Why should this be?" There may be physical, mental and/or social reasons, which then must be analysed in more detail to form the basis for decisions on rehabilitation.

"ADL assessment" is part of the broader concept of "functional assessment", which implies assessment not only of disability but also of impairments and handicaps. "Functional assessment" is to be found in DSM-III-R and in DSM-IV¹⁹ in Axis V. The Global Assessment of Functioning (GAF) scale is a hierarchical assessment scale not only of psychic symptoms but also of social and professional functioning.

The ADL instruments can be constructed as check lists, a summed index or hierarchical scales.

WHAT IS AVAILABLE TO ASSESS ACTIVITIES OF DAILY LIVING?

Many structured ADL assessment instruments have been developed but rather few have been used by persons other than their inventors. Most instruments include the same types of personal items, but they often have divergent operational definitions, if indeed they are defined at all. Some instruments combine personal and instrumental items. There are also multidimensional instruments with ADL items mixed up with those for other assessments, such as mobility, physical status, mental capacity and social conditions.

An overview of such instruments, mainly from Anglo-Saxon countries, is given in Kane and Kane⁴. This field of study is also undergoing continuous and rapid development in other countries, so readers are recommended to check up on the key-word "Activities of Daily Living" in *Index Medicus*.

DIAGNOSIS AND ASSESSMENT

When choosing an ADL instrument, three questions need to be asked:

Question 1: for what purpose is it to be used?

Possible uses could be:

- (a) To describe and document any need for assistance in performing ADL.
- (b) To differentiate among different levels of disability.
- (c) To follow and record changes in ADL performance over time.
- (d) To predict outcome, survival, length of hospital care, type and quality of living, etc.
- (e) To evaluate rehabilitation programmes, hospital, home and day care, etc.

Question 2: for whom is it intended?

- (a) For individuals, for groups with certain medical diagnoses or receiving different types of care or for populations?
- (b) For mainly healthy persons or for the mildly or severely disabled?
- (c) For patients living in institutions or in their own homes or for patients with or without cognitive impairment?

Most published instruments are shown to be reliable and valid for particular purposes and for particular patient categories. The same instruments may be unsuitable for use in other circumstances, e.g. instruments that include personal items only are often adequate for the more seriously disabled patients living in institutions, whereas instruments including instrumental items are necessary for use with more healthy populations living in their own homes. For patients with serious cognitive impairment, the ADL instrument has to depend on observation or test situations. The assessment has to take into consideration the patient's own dependence on active assistance as well as the patient's dependence on supervision, and whether or not activity is initiated by another person. The quality of ADL performance has also to be assessed; for example, if the patient dresses him/herself independently, does he/she dress him/herself adequately and appropriately for the situation?

Question 3: are the results of ADL assessment expected to be compared with other ADL assessments?

If so, it is necessary to choose an ADL instrument that has been tested and documented as to its reliability and validity for the chosen purpose and the specified patient category. If a group of patients is to be compared with normal subjects, data on nationally representative samples must also be available for comparison.

It may be difficult, if not impossible, to find one ADL instrument to fit all purposes. In any case, it is usually better to choose one of the well-known instruments and then supplement it with items that seem important in that specific situation, than to start the immense work of creating a new instrument. It is also better to use a plain ADL instrument instead of a multi-dimensional one, if the purpose is simply to study ADL ability.

Time and staff resources are also limiting factors when choosing an ADL-instrument. Self-administered questionnaires are less expensive to administer, but require patients who understand the questions and can respond adequately. The number of subjects who drop out may be high, so some validation of the answers is necessary. This type of instrument is mainly used in studies of elderly general populations, who are expected to be reasonably healthy.

Observations of the patient's actual ADL performance can be carried out during a nurse's daily work, when documentation will take only a few minutes. Training of the observers is usually required, along with tests of inter-observer reliability.

Test situations that use professional observers are the most expensive measures. Even if the test situation does not correspond exactly to real life, it can still be of value for specific purposes, e.g. when occupational therapists analyse the kind of impairment that is hindering performance or how rehabilitation should be undertaken; or in methodological studies, such as in a study of the relationship between severity of dementia and ADL performance⁵.

In clinical practice, occupational therapists usually combine self-report information concerning the disabled patient's own values, personal causation, interests, roles, habits and skills with observations and test situations⁶.

HOW GOOD ARE THE ADL ASSESSMENTS?

The quality of an ADL instrument depends on the following factors:

1. *Its representativeness:* i.e. whether the items reflect the activities of a person's normal daily life.
2. *Its reliability:* viz. its reproducibility and stability, so that the results can be reproduced by other observers or by the same observer at different times, so that changing results reflect changes in the patient's ADL status.
3. *Its validity:* i.e. whether the results are meaningful to others; whether they can be understood and used to compare with other data reflecting different levels of disability; and whether they may predict outcome.

These aspects of instrument quality are generally easy to test, but the problem is the lack of a norm or gold standard for comparison. The closest approximation to such a norm is the Katz Index of ADL⁷, which is based on a cumulative scale of personal items reflecting improvement and deterioration among disabled patients, and designed to correspond with ADL development in the small child.

Other instruments may include more or less the same items as in Katz Index, such as Barthel's Index⁸, but problems arise when the authors put arbitrary nominal values on different levels of ADL ability, which are ordinal in character, and then summarize the assessments. The results may be affected by systematic statistical errors, but more importantly they may obscure the understanding of actual ADL status and undermine the possibility for interstudy comparisons using different instruments. For example, if a patient scores 75 out of 100 possible points, this will not reveal in which items the patient is dependent. Furthermore, one patient with 75 points may not be as disabled as another person with an equal number.

Quite another way of handling ordinal data is shown by the development of Katz's Index of ADL. In the 1950s, a multi-disciplinary team in Cleveland, Ohio, followed patients who were recovering from a hip fracture by assessing in which order they regained the ability to perform ADL. They found six items that usually followed in sequence: first the patients regained independence in feeding, followed by continence, movement of the body (e.g. getting out of bed), going to the toilet, dressing and bathing. A patient who was independent in bathing could be expected to be independent in the other five items, too, and a person who was dependent in feeding was also dependent in the other items. The authors defined each item carefully, and found that dependence on another person, either by active assistance or supervision, decided whether or not the subject was dependent.

The authors had discovered a hierarchical or cumulative order between these six personal activities of daily life, which formed the beginning of an ADL "staircase" where each step upwards corresponded to an improvement in the patient's condition. Later, it was shown that patients with other diagnoses, such as stroke

and rheumatoid arthritis, and mentally impaired people improved and deteriorated in the same way⁹. The instrument has been used for a number of purposes among elderly and disabled people in many countries¹⁰. In Sweden, it is used in clinical practice by medical and geriatric departments and in home care to communicate information about the patients' ADL ability. The level of ADL performance has been shown to predict survival and death, length of hospital stay and type of hospital discharge in acute medical care^{11,12}.

The cumulative order of items was found to correspond to the observations made by Gesell of the ADL development in small children during their first 7 years. This sociobiological origin may explain the good predictive validity of the instrument. The limitation of the scale is primarily a ceiling effect. It does not differentiate among less disabled persons, who are independent in bathing. But the small child does not stop developing at age 7, and the very existence of a cumulative scale evoked the question of what comes after bathing.

From population studies, it was possible to derive scales of percentages of the population with activity limitations. Lawton and Brody¹³ formed hierarchical scales of personal and instrumental items. These cumulative ADL items were incorporated in a comprehensive multidimensional scale, the Older American Research and Service Center (OARS) instrument¹⁴. In order to provide shortened measures, Fillenbaum formed a five-item cumulative instrumental ADL scale out of the OARS instrument¹⁵. Spector and Katz¹⁶ reported that the instrumental items of cooking and shopping were ranked higher than personal items. From the Gothenburg population study¹⁷, ADL reduction was found in about 30% of 70 year-old people, mostly due to dependence in the personal items of the ADL.

To understand further the cause of disability in old age, it was necessary to formulate exact definitions for instrumental activities, which could be ordered into an extended cumulative scale. Sonn and Hulter Asberg¹⁸ found that the items of cooking, transportation, shopping and cleaning could be defined and ordered next to bathing in Katz's Index of ADL. The ADL "staircase" now consists of 10 items: feeding, continence, movement of the body (e.g. getting out of bed), going to the toilet, dressing, bathing, cooking, transportation, shopping, cleaning. This ADL staircase can be used for observation and documentation of the different levels of disability for individuals, groups of patients and for population studies. For individuals, it can be supplemented by test situations according to the occupational therapists' assessment of the need for rehabilitation. For groups of patients, it can be supplemented by disease- or symptom-specific scales, and populations can be assessed by interviews, where the interviewer also assesses whether or not the answer seems a reasonable one.

There may be special reasons for choosing other ADL instruments than the one described above, such as historical reasons, instruments that include more items and are more sensitive to small changes in disability and that offer the possibility of comparison with earlier studies. Furthermore, important research work by occupational therapists is now in progress, attempting to develop special ADL instruments for

patients with psychiatric problems. The field is, as it were, "preadolescent". In order to achieve a general standard for ADL assessments, it would be valuable to use the ADL staircase alongside other instruments. This would facilitate comparisons between different studies and help promote knowledge about, and understanding of, the ADL status of elderly patients.

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Rating Scales Designed for Nurses and Other Workers

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The changing role of nurses and other workers in the context of clinical audit and community care has provided a fertile backdrop for the development of brief and easy-to-use clinical rating instruments. Few of these are of diagnostic validity and they require little or no training. They are usually employed to rate clusters of signs, symptoms, syndromes, behaviours or needs. They have often been developed from other instruments or through clinical observation. The majority have been exposed to validation and inter-rater reliability and some are furnished with “cut-off” scores corresponding to external diagnostic criteria. Nurses and other professionals have been successfully trained in the delivery of more complex diagnostic and assessment instruments, including CAMDEX and GMS. However, this chapter will focus on relatively brief clinical instruments requiring little or no training, primarily designed for nurses and carers. This chapter is by no means exhaustive but is designed to provide an overview in a rapidly developing field.

ORGANIC STATES INCLUDING DEMENTIA

Some of the specialized neuropsychological instruments, including the Kendrick Battery, are usually carried out by psychologists and have been extensively validated¹. However, there now exists a wide range of brief cognitive assessments that have gained both scientific and clinical credibility, readily available to clinicians of all professions. The Mini-Mental State Examination² is pre-eminent in this field. It is easily administered and requires no training. The test consists of verbal and performance components. The verbal subtests evaluate orientation in time, memory and attention. The performance subtests involve the naming of objects, execution of written and spoken orders, writing, and copying a complex polygon. It takes a relatively short time to administer, and can be carried out by medical and paramedical staff. It has been extensively validated and has high inter-rater and test-retest reliability. Culture and social status significantly affect the scores³ but it has been employed in the context of transcultural epidemiological studies in which modifications have been made⁴. The Standardized Mini-Mental State Examination is a derivative designed to promote standardization and inter-rater reliability⁵.

The Blessed Dementia Rating Scale⁶ has been a source scale for the development of further rating instruments. Cognitive assessment includes memory and information items. The second part consists of items of behaviour. The scale has been validated through correlation with mean brain cortical plaque counts. It “ceilinged” with the correlation declining sharply in clinically

severe dementia sufferers. The 10 most useful questions differentiating between normal and abnormal cognitive functioning have been extracted⁷ and are used as the Abbreviated Mental Test Score, giving comparable results to the full Blessed scale. The latter has been used in a simple test of mental impairment in functionally and mildly organically ill patients attending a day hospital⁸ and has been used in screening for organic disorder and predicting change over a 2-year period⁹. The study evaluated the degree of disability and was used in the planning of social support.

The Clifton Assessment Procedures for the Elderly (CAPE) consists of two sub-scales, the Cognitive Assessment Scale and the Behaviour Rating Scale, derived from the Stockton Geriatric Rating Scale, designed to be used by nurses¹⁰. It was devised as a brief measure of psychological functioning for chronic psychiatric patient groups and has been validated against the outcome of day hospital and day centre care¹³. There is poor inter-rater reliability of the Gibson Maze component, which has to be completed by the patient. Despite these problems, the CAPE provides a well-validated, useful instrument that does not take too long to complete. Other common instruments include the Mental Status Questionnaire¹⁴ (MSQ), requiring some training, as questions are asked in a standardized fashion. The Short Orientation–Memory–Concentration Test was developed from the MSQ and consists of six items concerning orientation and memory¹⁵. One of the more recent developments has been the Clock Drawing Test, which assesses frontal and temporoparietal functions, providing a useful bedside assessment¹⁶.

SCALES FOR RATING BEHAVIOUR AND SELF-CARE

The Stockton Rating Scale (from which items of the CAPE were derived) is pre-dated by the Crichton Behavioural Rating Scale (CBRS)¹⁷. The CBRS rates 10 aspects of behaviour, including the functions of orientation, communication and mood. The addition of a memory item allows the generation of a “confusion” score that is sensitive to change in mentally ill patients. A modified version has been used in the context of nursing home residents.

The Performance Test for Activities of Daily Living (PADL)¹⁸ was developed for the US/UK Cross-National Project. It assesses the degree of autonomy of elderly psychiatric subjects in activities of daily living. It is presented as a test of praxis containing 16 subtests. Lawton and Brodie developed the Instrumental Activities of Daily Living Scale (IADL) and the Physical Self-maintenance Scale¹⁹. The IADL is widely used to evaluate the degree of

physical and instrumental autonomy of elderly subjects. It takes 5 min to deliver and has been used as a base instrument for more recent developments in this field. Both the IADL and the PADL are task-based scales using simple scoring systems.

The Stockton and other scales²⁰ inspired the Physical and Mental Impairment of Function Evaluation in the Elderly (PAMIE). The assessment is completed with the aid of simply formulated items not requiring rater interpretation. It measures behavioural characteristics in patients suffering from chronic diseases, with particular reference to the elderly. Two other scales, the Psychogeriatric Dependency Rating Scale (PGDRS)²¹ and the Geriatric Behavioural Scale²², have been developed by nurses, psychologists and psychiatrists. Both scales have been used for the assessment of dementia and physical incapacity. They have been well validated in the context of prognostic ability over a period of 1–2 years. The only scale designed to examine behaviour in elderly schizophrenic subjects is the Nurses' Observation Scale for Inpatient Evaluation (NOSIE)²³. The scale consists of 30 items and takes about 20 min to complete. It is based on 3-day consecutive observation of the patient and has a frequency scale of 0–4 for each item.

A number of scales designed to be completed by nurses and carers address disruptive, antisocial and aggressive behaviour in dementia sufferers. These include the Overt Aggression Scale²⁴, The Disruptive Behaviour Rating Scale²⁵, The Nursing Home Behaviour Problem Scale²⁶ and the Brief Agitation Rating Scale²⁷. The Caretaker Obstreperous-Behaviour Rating Assessment (COBRA)²⁸ and the Ryden Aggression Scale²⁹ are slightly longer, taking up to 30 min to complete.

SCALES ASSESSING CARERS AND ENVIRONMENT

The Multiphasic Environment Assessment Procedure³⁰ is a relatively well-validated questionnaire for the assessment of the environment in which dementia sufferers are nursed. It enables the assessment of the complex relationships existing between physical environment, the policies and the characteristics of the staff and residents. Adaptations have to be made for its use in the UK, as it was designed and validated in North America³¹. The Social Network Assessment Scale³² has been used in typing social networks of older dementia sufferers.

SCALES ASSESSING CARERS

Gilleard³³ developed the Problem Checklist and Strain Scale, designed to assess problems experienced by carers. Other less well-known instruments include those designed to assess psychological problems facing the carer: the Ways of Coping Check List³⁴ and the Burden Interview³⁵. The Relatives' Stress Scale³⁶ enables the relatives to make a standard assessment of stress they are experiencing as a result of having to care for an elderly demented person living at home. The Caregiver Activity Survey³⁷ quantifies the time a caregiver devotes to the patient, and the Marital Intimacy Scale³⁸ specifically examines the psychological and emotional consequences of caring for an ill spouse.

RATING SCALES FOR MORALE AND MOOD

The Measurement of Morale in the Elderly Scale consists of items extracted from empirical observations³⁹. It is a long, structured interview lasting 2–4 h. A degree of training and knowledge of the instrument is required before it is used, in order to achieve satisfactory levels of reliability. A shorter morale scale, The Philadelphia Geriatric Centre Morale Scale⁴⁰, takes 10 min and

requires no training. The questions are of a forced-choice type and are read to the respondent. There is an internal consistency of factors and the scale has had fairly wide use. It is a multi-dimensional instrument for use in the very old and was particularly designed not to provoke fatigue or excessive inattention.

Depression rating scales should only be employed in rating depression in older people if validation studies have been reported. Some rating scales have been specifically designed for this age group, accommodating the issues of concomitant physical illness and cognitive change. The Self Care (D)⁴¹ is a self-administered scale. It has been used in the community for the screening and rating of severity of depression in the elderly. It was found to be a sensitive indicator of depression. It may have potential use in the monitoring of change as a consequence of antidepressant medication. The scale has been validated in a pilot study in elderly, continuing-care patients⁴². The Geriatric Depression Scale (GDS)⁴³ was devised by compiling 100 questions used by professionals in the diagnosis of depression. O'Riordan *et al.*⁴⁴ recently used the scale in a large survey of patients admitted to an acute medical geriatric assessment unit, where there was a high prevalence of physical illness and cognitive impairment. They found that the GDS appeared to be a sensitive test for depression in this group, but was sufficiently non-specific to require psychiatric evaluation of patients with high scores to establish the accurate prevalence of depressive illness. The instrument exists in 15-item⁴⁵, 10-item and four-item versions⁴⁶. Other scales that are used by clinicians include the Cornell Scale for depression in dementia⁴⁷, the Brief Assessment Schedule Depression Cards⁴⁸ and the ELDRS⁴⁹ designed to be used with an informant as well as the patient. The Centre for Epidemiological Studies Depression Scale (CES-D)⁵⁰ is a self-rating scale that has been found to be of value in screening community samples.

CONCLUDING REMARKS

This chapter does not provide a comprehensive account of the wide variety of instruments that are available for the assessment of the elderly mentally ill. Attention has been given to those instruments that have been relatively well validated, with evidence of reliability, that are easy to use with little or no training. They have been loosely categorized by function and use. Care must be taken in their application, with appropriate supervision and standardization in view of frequent problems of inter-rater reliability. In isolation, these instruments are of limited clinical value, but in the context of a multi-professional team they do have important potential in screening, clinical audit, examining issues of service need and rating change.

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Comprehensive Interviews

OARS Methodology

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The OARS (Older Americans Resources and Services) methodology was developed over two decades ago in response to a request “to assess alternative strategies to institutionalizing frail older adults”¹. To address this issue, a three-element model was developed, which allows one to: (a) assess adults at all levels of functioning from excellent to totally impaired, using a multidimensional perspective; (b) determine the extent of use of and perceived need for each of 24 broadly encompassing but non-overlapping generically-defined services, which can be aggregated into costable packages; and consequently (c) to examine the impact of the identified service packages on persons defined in terms of their multidimensional functional profiles.

The questionnaire developed to operationalize the first two elements of the model is not restricted to it, but has enjoyed more varied use. The questionnaire is in two parts (see summary in Table 1). Part A assesses the level of functioning in five areas. Three areas reflect personal functioning—mental health, physical health, and activities of daily living; two reflect environmental conditions—social resources and economic resources. In each area the information obtained (from the subject or an informant) can be summarized on a six-point scale, ranging from excellent to totally impaired functioning. Review across all five areas provides a profile, making it possible to identify where functional strengths and weaknesses lie. Each area is itself multidimensional, so allowing users to examine specific aspects of particular areas of functioning and permitting more accurate identification of service impact.

Part B focuses on services assessment. To ensure accurate identification, each service is defined in terms of its purpose, the activity involved, the personnel who may provide the service, and the units in which it is to be measured (to facilitate costing). So, a resident of a nursing home would not be rated as receiving “nursing home services”, rather, the precise services received there (e.g. nursing care, meal preparation, occupational therapy), the amount, and the type of provider (formal or informal) would be recorded. Need for services is self-assessed.

On average, the OARS questionnaire takes 40 min to administer (training in this is provided). Validity and reliability have been determined. It is available, and has been validated, in a number of languages. The questionnaire has been used for purposes as varied as teaching, clinical and population assessment, agency evaluation, staffing determination, service impact, prediction of service needs, determination of preferred service aggregation, and estimating service cost in different settings. An

Table 1. Overview of OARS multidimensional functional assessment questionnaire

Part A, Assessment of functional status	Part B, Services assessment
Demographic	Transportation
Social resources	Social/recreational services
Interaction	Employment services
Affect	Sheltered employment
Extent of availability of help	Educational services, employment-related
Economic resources	Remedial training
Occupation	Mental health services
Income (by source)	Psychotropic drugs
Housing	Personal care services
Self-assessed adequacy of income	Nursing care
Mental health	Medical services
Short Portable Mental Status Questionnaire (to assess level of cognitive functioning)	Supportive devices and prostheses
Short Psychiatric Evaluation Schedule (to assess presence of psychiatric problems)	Physical therapy
Self-assessed mental health	Continuous supervision
Physical health	Checking services
Prescribed medications	Relocation and placement services
Current illnesses and conditions and their impact	Homemaker–household services
Alcohol use	Meal preparation
Level of activity	Administrative, legal and protective services
Self-assessed physical health	Systematic multidimensional evaluation
Activities of daily living (ADL)	Financial assistance
Instrumental ADL	Food, groceries
Physical ADL	Living quarters (housing)
	Coordination, information and referral services

archive of OARS-based data sets is maintained at the Duke Aging Center.

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The Comprehensive Assessment and Referral Evaluation (CARE): An Approach to Evaluating Potential for Achieving Quality of Life

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PURPOSES

The CARE covers a wide range of indicators of a person's potential for achieving a preferred quality of life. Its bounded focus is on health and social problems associated with aging, including psychiatric disorders. Originally developed for intensive studies of community-dwelling elders¹, the rater-administered CARE has given rise to versions that have been made briefer by concentrating on key scales (the CORE-CARE² and the SHORT-CARE)³, suitable for residents of institutions (the IN-CARE)⁴, clinically focused (the CLIN-CARE)⁵, and capable of self-administration (the SELF-CARE). There is also an INFORMANT version⁶ of the CARE.

THEORETICAL BASE⁷

The phrase "quality of life" is invoked to direct attention to processes of striving to achieve or preserve a preferred manner of living. The processes involve strategies that allow relevant choices to be made and implemented. The strategies can be impeded by certain health and social problems that may occasion help-seeking contacts with informal or formal caregivers or services. The CARE seeks to assess the person's ability to deal with the strategies necessary for choosing and attaining a preferred manner of living. It does not attempt to define the latter. Information on processes can come from many sources, touching on many aspects of living, reaching beyond the person's immediate status into the present and historical context, modified by expectations of the future, and qualified by personal and cultural preferences. It is recognized that the gathering of such information is very constrained by its complexity, openness to change and inherent uncertainties, as well as by the necessary rigidities of the systematic assessment procedures.

STYLE

The general style of the CARE relies on scripted questions with pre-coded answers. The questioning is tactful and with an organization that is understandable to the interviewee. Header questions allow sections to be skipped if unlikely to be productive. Computer-assisted programs guide interview administration, so as to avoid missed or conflicting ratings. Information can be analyzed at the level of discrete items, global ratings, scales, hierarchically-arranged classes of severity, and threshold scores relating to diagnosis and need for investigation and possible treatment.

DOMAINS

Each domain targets the capacity to meet a distinctive challenge to achieving a preferred quality of life. Nineteen quality of life

domains⁸ have been identified and matched to items in the CARE: moving purposefully (mobility); maintaining self routinely (basic activities of daily living); using the immediate environment (instrumental activities of daily living); manipulating household appliances (technological activities of daily living); finding one's way around (navigational orientation); keeping track of time and space (continuity orientation); gathering information (receptive communication); expressing needs (expressive communication); preserving health (health and safety practices); protecting physical and mental comfort (emotional and physical status); engaging in social relations; exercising choice; managing material resources; finding the best environment; obtaining meaningful gratifications; recognizing one's own state of health (self-perceived health); taking account of the future (pessimism, optimism, realism); balancing competing qualities of life; setting and achieving goals. The domain content and labels are keyed to adaptive behaviors in pursuit of quality of life (more conventional captions are shown in parentheses).

SUBJECTIVITY AND OBJECTIVITY

The CARE does not beg the issue of whether quality of life is primarily subjective or objective, preferring to regard both as important strategies in the pursuit of quality of life. Subjective (e.g. feelings, attitudes), quasi-objective (self-reports of objective status) and mainly objective (tests and observations) aspects of quality of life are represented, thus allowing their interrelationships and their respective effects and outcomes to be examined. For example, questions on activities of daily living probe self-reported task performance, views on the extent to which health limits desired activities, informant views on the elder's functioning, tests of range of movement and observations on mobility. A supplement tests a simulation of various basic and instrumental tasks (the Performance of Activities of Daily Living, or PADL)². Similarly, inquiries on cognitive status range from self-reported difficulties with memory to formal tests of memory and orientation, and a supplement (the Medication Management Test, or MMT)^{10,11} tests higher-order cognitively-driven performance.

SEVERITY LEVELS

Threats to quality of life are graded in terms of the degree to which they overshadow living at a critical point in time (intensity) and over time, experiences, and situations (extensity). This approach has been more completely modeled within the CARE domain of emotional comfort, as expressed in the seven levels of the index of affective suffering (IAS)^{12,13}. Each level is operationalized by symptom criteria that convey the severity meanings of suffering in a way not equaled by symptom scores or diagnosis. This model of severity is also worked out for the CARE domains dealing with functioning in the activities of daily living. Current development is concerned with deriving a measure

of the continuities underlying the progression of functional decline.

MECHANISMS

Threats to quality of life, such as those measured by the CARE, may arise from health and social conditions; the presence of the latter can be suggested by certain syndromes. The CARE items include syndromes of cardiac failure or angina, arthritic pain, respiratory problems, effort intolerance, cognitive deficits, Parkinson's disease, stroke, perceptual deficits, vertigo, falls, experience of crime, and social isolation and desolation. Conversely, there are characteristics of the threatened qualities of life that suggest certain mechanisms: one scale of functioning in the CARE is composed of higher-order tasks that are principally affected by cognitive mechanisms; similarly, certain somatic items are phrased to suggest a mood disorder and others to suggest a physical disorder. More directly, there are items to record what the individual blames as the cause of his/her threatened quality of life.

BIOMETRICS

Satisfactory reliability, validity and operational characteristics for discriminating diagnoses of the CARE scales have been established¹⁴⁻¹⁸. Transition tables show a range of incidence, chronicity and recovery from threats to quality of life¹⁹; these also reflect a power to predict specific quality-of-life outcomes. Cross-tabulations reveal that the various domains are sufficiently independent to be usefully measured separately but that there is substantial interaction between them.^{20,21}

CULTURE-FAIR INDICATORS

Ethno-racial and educational variation in qualities of life have been found with CARE indicators^{22,23}, as with comparable instruments²⁴. In order to minimize the confound of bias entering into such comparisons, item response theory has been applied to construct culture-fair CARE scales of mood, cognition and functioning in the activities of daily living²⁵.

MODULAR USES

Although the full CARE is more than the sum of its parts, the scales can be, and have been, used alone and in various combinations. This has enabled the assembly of its different versions.

WHOLE-PERSON VIEW²⁶

Profiles of CARE indicator severity scales are generated by computer algorithms, giving an overview of the person's strengths and vulnerabilities for preserving and improving quality of life. Global concepts are also embodied in items on self-perceived general health and in visual analog scales on physical and psychological distress and task and social functioning: a supplementary single-page rapidly completed set of visual analog scales (the QoL-100) has proved useful for a snap-shot of the quality of life potentials of seriously ill patients or where frequent follow-up assessments are needed. Psychiatric diagnoses have been generated using a computerized decision tree^{27,28}.

APPLICATIONS

Taken together with its versions and supplements, the CARE has been applied to epidemiological²⁹ surveys³⁰⁻³³, public health policy³⁴⁻³⁹ and clinical management in primary medical care⁴⁰, home care⁴¹ and occupational therapy. A Training Manual is available.

FURTHER DEVELOPMENT

The Stroud Center's program is designed to contribute to understanding the nature of the parts and the connected whole of a person's quality of life. Experience with applications, modifications and analyses of the CARE is leading to new and better ways of assessing quality of life.

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Part E

Organic Disorders

- EI Delirium
- EII Dementia
- EIII Alzheimer's Disease
- EIV Vascular Dementia
- EV Other Dementias
- EVI Clinical Diagnosis of the Dementias
- EVII Outcome of the Dementias and Subtypes
- EVIII Treatment and Management of Dementias
- EIX Conditions Associated with, or Sometimes Mistaken for, Primary Psychiatric Conditions
- EX Investigation of Organic States and Dementia

Delirium—An Overview

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Delirium is a state of acute confusion due to an underlying physical cause. Associated with the disorder of cognition and attention, there is frequently disturbed psychomotor behaviour and disturbance to the sleep-wake cycle. It is a common feature in those, particularly older patients, who present as acute medical emergencies. As a common, potentially highly treatable condition, it is associated with major health care costs. It is also under-recognized and associated with longer periods of inpatient stay. It is a recognized mental disorder under the English Mental Health Act, which has significance in relation to compulsory treatment.

Formal diagnostic criteria for delirium were only first introduced in DSM-III. These criteria were further revised for DSM-IV and the criteria for general medical conditions are shown in Table 35.1. DSM-IV criteria are also described as more specific for delirium in relation to substance abuse. The main changes in the criteria are a general simplification with less emphasis on deficits of attention and more emphasis on the syndrome developing over a short period of time.

Confusion, or cognitive impairment, commonly only occurs in three conditions, namely delirium, dementia and depression. Two of the “3Ds” are eminently treatable, therefore it is to be regretted that cognitive impairment is primarily associated with dementia and hence only belatedly recognized as a potentially treatable condition in the case of this particular chapter’s subject, delirium.

The rest of the chapter will discuss incidence and risk factors. There is an expanded section on clinical features and discussion of assessment scales, neuropathogenesis, appropriate investigations and treatment.

INCIDENCE

Delirium occurs in 14–56% of older hospitalized patients¹. It is lower after elective surgery and higher in acute medical admissions. Patients who have suffered delirium during a hospital admission tend to stay in hospital for longer and have greater requirements for rehabilitation and require increased home care services².

RISK FACTORS

Inouye *et al.*¹ propose a multifactorial model for delirium, including predisposing factors and precipitating factors. The study identifies four predisposing factors for delirium, visual impairment, severe physical illness, cognitive impairment and blood urea nitrogen (BUN):creatinine ratio i.e. an indicator of dehydration. Using complex methodology, they identified a

hierarchy of precipitating factors, the most significant of which were major surgery, stay in intensive care, multiple medication and sleep deprivation. Their conclusion was that the aetiology of delirium is multifactorial, but that certain predisposing factors, combined with a weighting of precipitating factors can make the likelihood of delirium much more predictable. Elie *et al.*³ similarly identified risk factors that included dementia, advanced age and medical illness. Robertsson *et al.*⁴ looked at the likelihood that a pre-existing dementia could predispose to the onset of delirium and showed that late-onset Alzheimer’s disease was more likely to do so than early-onset Alzheimer’s disease and that vascular dementia was more likely to do so than early-onset Alzheimer’s disease.

CLINICAL FEATURES

Whilst DSM-IV attempts to define the clinical features, it is worth remembering that Lipowski⁵ has regularly warned about the subtle “prodromal, non-specific” features that can even precede the more formal syndrome. This can be as simple as increased anxiety or subtle levels of agitation. The cardinal feature is probably a relatively sudden change in cognition but there can also be perceptual disturbances. Patients can latterly describe great difficulties in differentiating between reality and hallucination. Visual hallucinations are classically more common than auditory hallucinations and are generally unpleasant and frightening. Thinking is disorganized, slowed and impoverished. Memory is impaired across the entire spectrum of registration, retention and recall. Short-term memory may well be impaired, secondary to poor attention, and patients may be inclined to confabulate.

The impact of delirium on psychomotor behaviour is variable. Some delirious patients will become agitated, restless and hypervigilant. In contrast, some patients will become withdrawn, with slowed physical responses. Finally, the picture of psychomotor activity can be mixed with oscillation between hyper- and hypo-activity.

Having described the classic clinical features above, it should be remembered that Treloar and McDonald⁶ have placed emphasis on the “quiet syndrome”. They argue that one of the reasons that delirium is under-recognized is because of the frequency of the hypo-active type of syndrome. There is evidence that nurses are better than doctors at identifying the syndrome of delirium and there is tentative evidence that different aetiologies may possibly account for differentiation in the clinical picture. Zou *et al.*⁷ showed that diagnosis by a nurse clinician using a standardized rating scale (the Confusion Assessment method) and multiple

Table 35.1 Diagnostic criteria for 293.0 Delirium, due to... (indicate the general medical condition)

-
- A. Disturbance of consciousness (i.e. reduced clarity of awareness of the environment) with reduced ability to focus, sustain, or shift attention
 - B. A change in cognition (such as memory deficit, disorientation, language disturbance) or the development of a perceptual disturbance that is not better accounted for by a pre-existing, established or evolving dementia
 - C. The disturbance develops over a short period of time (usually hours to days) and tends to fluctuate during the course of the day
 - D. There is evidence from the history, physical examination or laboratory findings that the disturbance is caused by the direct physiological consequences of a general medical condition
-

Coding note: If delirium is superimposed on a pre-existing Dementia of the Alzheimer's Type or Vascular Dementia, indicate the delirium by coding the appropriate subtype of the dementia, e.g. 290.3 Dementia of the Alzheimer's Type, with Late Onset, with Delirium.

Coding note: Include the name of the general medical condition on Axis I, e.g. 293.0 Delirium Due to Hepatic Encephalopathy; also code the general medical condition on Axis III (see Appendix G for codes)

observation points was better than a one-off assessment by a psychiatrist. This study reinforces the importance of repeated observations and the value of a structured approach to the diagnosis.

ASSESSMENT SCALES

Because of the problem clinicians have in recognizing delirium, there have been a number of attempts to create assessment scales to assist in diagnosis. In the development of these scales, one of the major problems has been that cognitive impairment is, of course, a common symptom but that it is not specific to delirium. A number of instruments to evaluate delirium have been developed, mainly using DSM-III-R criteria. Many of them are multi-item scales, which probably have limited value in the clinical setting, but one, the Confusion Assessment Method (CAM), is intended to be used by physicians and has two versions with only nine and four items, respectively. Non-specific cognitive tests, such as the well known Folstein Mini-Mental State Examination, cannot specifically address the differentiation between delirium and dementia. However, it would seem that the introduction of appropriately clinically friendly scales for delirium, used by skilled nurse clinicians, might well increase the recognition of delirium in acute medical settings⁷.

NEUROPATHOGENESIS

Whilst delirium is regarded as a syndrome with global dysfunction of cognition, attempts have been made to address whether there are disorders of specific brain pathways that can account for specific symptomatology, the analogy being that certain specific strokes can be associated with syndromes, for example, left frontal strokes being associated with depression. Trzepacz⁸ has reviewed a number of brain scan studies which, although they are limited to studies with only a small series or even single case reports, tend to show that frontal cortex, anteromedial thalamus, right basal ganglia, right posterior parietal cortex and medial basal temporal and occipital cortex may be areas particularly associated with delirium.

The neurochemistry of delirium is believed to mainly involve under-activity of the cholinergic system. That said, on the basis that most neurochemical systems in the brain are in dynamic relationships with others, it is highly likely that this acetylcholine hypothesis also relates to a balance with the dopamine system. It can be speculated that, in the future, there may be scope to manipulate the balance of the acetylcholine and dopamine systems in such a way as to treat the symptoms of delirium, very possibly with the new cholinesterase inhibitors.

INVESTIGATIONS

Delirium, particularly in the elderly, tends to have a multi-factorial aetiology. As it can be due to almost any underlying physical problems, the old adage that "common things occur commonly" must be relevant when investigating the causes. These are likely to be infections, cardiac problems, iatrogenic medication, dehydration, stroke disease, diabetes and cancer. Naughton *et al.*⁹ reported a study of head CT scans in delirium. 15% were positive for an acute condition (haemorrhage, haematoma, space-occupying lesion or infarct) and 95% of these positive scans were found in patients with impaired consciousness or a new focal neurological finding detected during the episode of delirium. Their main conclusion was that there was considerable variation in brain CT scanning of older people with delirium.

TREATMENT

First, the aetiological factors must be identified and treated or corrected if at all possible. Whilst these are being identified, symptomatic and supportive therapies will be required. Over the years, Lipowski⁵ in particular has recommended an appropriate environment for managing patients with delirium. He has described a well-lit room with familiar items and clearly visible clock and calendar. Inouye *et al.*¹⁰ have systematically studied this in a paper describing a multi-component intervention to prevent delirium in hospitalized older patients. Their risk factors and intervention protocols are detailed in Table 35.2. The study provides important evidence that a multicomponent, targeted intervention to prevent delirium in hospitalized older patients can work, particularly targeting those at high risk from previously identified precipitating and predisposing factors. Wahlund and Björlin¹¹ report on a specialized delirium ward and argue that such a specific ward is best placed to clinically manage, investigate and treat patients with delirium. Despite the importance of primary prevention and the probable value of special delirium wards, there may be a need for antipsychotic medication. Haloperidol probably remains the antipsychotic of choice¹², although little is currently known about the efficacy of the new atypical antipsychotics in delirium. Benzodiazepines probably remain the preferred medical treatment in delirium due to alcohol.

As noted above, delirium *per se* makes a longer hospital stay likely, with consequent increased health care costs. That said, the mean duration of delirium is shorter in post-operative patients compared to acute medical patients¹³. The long-term prognosis of delirium used to be felt to be simply that of the underlying condition. However, a recent publication¹⁴ flags up the possibility that the long-term prognosis after delirium is poor and, indeed, delirium might be a marker for decline. Also in contrast to traditional views, there is now a view that delirium is not always entirely reversible.

Table 35.2 Risk factors for delirium and intervention protocols

Targeted risk factor and eligible patients	Standardized intervention protocols	Targeted outcome for reassessment
Cognitive impairment* All patients, protocol once daily; patients with base-line MMSE score of 520 or orientation score of 58, protocol three times daily	Orientation protocol: board with names of care-team members and day's schedule; communication to reorientate to surroundings Therapeutic activities protocol: cognitively stimulating activities three times daily (e.g. discussion or current events, structured reminiscence or word games)	Change in orientation score
Sleep deprivation All patients; need for protocol assessed once daily	Non-pharmacologic sleep protocol: at bedtime, warm drink (milk or herbal tea), relaxation tapes or music, and back massage Sleep enhancement protocol: unit-wide noise reduction strategies (e.g. silent pill crushers, vibrating beepers and quiet hallways) and schedule adjustments to allow sleep (e.g. rescheduling of medications and procedures)	Change in rate of use of sedative drug for sleep [†]
Immobility All patients; ambulation whenever possible, and range-of-motion exercises when patients chronically non-ambulatory, bed- or wheelchair-bound, immobilized (e.g. because of extremity fracture or deep venous thrombosis), or when prescribed bed rest	Early mobilization protocol: ambulation or active range of motion exercises three times daily; minimal use of immobilizing equipment (e.g. bladder catheters or physical restraints)	Change in Activities of Daily Living score
Visual impairment Patients with 520/70 visual acuity on binocular near-vision testing	Vision protocol: visual aids (e.g. glasses or magnifying lenses) and adaptive equipment (e.g. large illuminated telephone keypads, large-print books and fluorescent tape on call bell), with daily reinforcement of their use	Early correction of vision, 448 hours after admission
Hearing impairment Patients hearing 46 of 12 whispers on Whisper Test	Hearing protocol: portable amplifying devices, earwax disimpaction, and special communication techniques, with daily reinforcement of these adaptations	Change in Whisper Test score
Dehydration Patients with BUN:creatinine ratio 418, screened for protocol by geriatric nurse-specialist	Dehydration protocol: early recognition of dehydration and volume repletion (i.e. encouragement of oral intake of fluids)	Change in BUN:creatinine ratio

*The orientation score consisted of results on the first 10 items on the Mini-Mental State Examination (MMSE).

[†]Sedative drugs included standard hypnotic agents, benzodiazepine and antihistamines, used as needed for sleep.

Finally, delirium is a mental disorder and as such comes under the English Mental Health Act. Very occasionally, therefore, it may be legitimate to use the Mental Health Act to facilitate the treatment of delirium.

SUMMARY

Delirium remains a common, under-diagnosed symptom of underlying physical illness in older people. This alone makes it an important health economic issue. Primary prevention of delirium is now of proved efficacy and, generally, the prognosis of delirium is that of the underlying physical disorder.

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Delirium in Institutions

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The prevalence of delirium is approximately 10–40% of patients on acute medical and surgical units. The incidence for newly admitted patients ranges from 25% to 60%¹. Delirium is commonly misdiagnosed as depression; up to 41.8% of inpatient psychiatry consultations for depression actually result in a diagnosis of delirium^{2,3}. Delirium has a number of adverse outcomes, and may itself be looked upon as a “symptom” of problems in the delivery of hospital services⁸. It is the single most important factor contributing to in-hospital complications, such as falls and pressure sores⁴. It has a significant impact on increasing hospital length of stay^{4,5} and upon the need for discharge to long-term care institutions⁶. It is the single strongest predictor of impaired daily function (AFL) at 6 months⁷, although most studies have found surprising little impact on mortality^{4,7}. Delirium may not be as reversible as previously thought; Levkoff found that only 17.6% of all new symptoms of delirium had cleared fully at 6 month follow-up⁵. Delirium is the most frequent complication of hospitalization in older patients.

DIAGNOSIS

Careful diagnosis of delirium is essential, as up to 66% of cases are missed¹. Delirium can be systematically diagnosed using the Folstein Mini-Mental State Examination (MMSE)⁹ in combination with the Confusion Assessment Method¹, with the optimal technique using several observation points¹⁰. Inattention is a critical criterion of delirium and is essential in differentiating it from depression and dementia. A recent model developed by Inouye and colleagues¹ at the Yale University Elder Life Program demonstrates that the probability of developing delirium is somewhat predictable among a population of vulnerable patients characterized by older age, high medical co-morbidity, sensory (visual/hearing) impairment and baseline cognitive impairment. Among this vulnerable group, the introduction of specific precipitating causes of delirium (such as indwelling catheter, use of restraints, and new medical complications) has been shown to increase the probability of delirium in proportion to the number of precipitating factors present.

TREATMENT

As yet, despite promising evidence that neuroleptics may improve the course of delirium itself¹¹ and that cholinesterase inhibitors may be effective¹², there are as yet no specific recommended pharmacological treatments for delirium. However, there is more

hopeful evidence that delirium may be prevented. An intervention based on Inouye’s approach provides: sensory correction aids (e.g. glasses and hearing aids); reorientation; therapeutic cognitive activities; a non-pharmacological sleep protocol; a dehydration protocol; and an early-mobilization protocol. This resulted in a significant reduction in new cases of delirium among patients at intermediate baseline risk for delirium. It is notable that this strategy had no impact on “delirium in progress”; its primary clinical impact lay in prevention¹³. Further work on interventions for delirium will clearly be of vital importance to the reduction of the risks for delirium in institutions and for its treatment. The costs incurred by early detection may well be more than offset by savings in decreased length of stay, decreased rates of institutionalization and decreased rates of in-hospital complications⁸.

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Prognosis of Delirium

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Studies of outcome have started to challenge the assumption that delirium is a truly reversible disorder with a good prognosis. Prospective outcome studies of delirium are required to describe its prognosis. Using change in MMSE¹ score as the outcome measure, Fields *et al.*² found that by no means all patients improve, and some get worse. Treloar and Macdonald³ found that although many acute patients had reversible cognitive dysfunction, few regained normal cognition. Some patients took the full 3 month time-course of the study to start their recoveries. Studies based upon standard diagnoses of delirium have also reported poor recovery at follow-up⁴. Kolbeinsson and Jonsson⁵ found that elderly patients with DSM-III-R⁶ delirium but not dementia at outset, had dementia at follow-up in 70% of cases. Cole *et al.*⁷ found that less than 50% recovered mentally at follow-up. Rudberg *et al.*⁸ showed delirium lasting over periods of several weeks (even though patients did not meet Delirium Rating Scale⁹ criteria for delirium continuously). Cognitive deficits and behavioural abnormalities persisted in the majority of patients for at least 6 months¹⁰ Kaponen *et al.*¹¹ found infrequent good cognitive outcome following delirium at one year.

Delirious patients stay longer in hospital than those without delirium. Diagnosis-related group length of stay of 13 days for patients with delirium compares with 3.3 days for those with dementia¹². Physical morbidity may be exacerbated by psychiatric co-morbidity. Delirium is associated with death in up to 33%¹³. It is frequently asserted that mortality is purely the result of the physical illness, but this is a difficult hypothesis to test. It would be very surprising if the stupor, retardation and poor compliance with treatments seen in delirious patients did not contribute to mortality from physical illness. Patients with delirium after hip fracture surgery are at greater risk of incontinence, urinary tract infections and pressure sores¹⁴. Delirium in patients with hip fractures also predicts higher long-term mortality and poor functional recovery¹⁴⁻¹⁶. Francis and Kapoor¹⁷ found that delirium predicted a doubling of mortality attributable to functional impairment, and in survivors predicted loss of ability to live independently. Levkoff *et al.*¹⁰ found that new symptoms of delirium resolve at 6 months in only 17.7%.

The prognosis of delirium is almost certainly not, therefore, one of early full recovery. Rather, delirium is a condition with a slow recovery and one which often fails to resolve completely.

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Nosology of Dementia

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In the early 1950s, Sir Martin Roth defined dementia as “severe decline in memory accompanied by disorientation for time and place”. The modern criteria emphasize that dementia is a global decline of intellectual functions that affects more areas than just memory. However, memory impairment is mandatory for the diagnosis in the *Diagnostic and Statistical Manual of Mental Disorders*, Version III—Revised (DSM-III-R) and IV (DSM-IV) (see Tables 36.1 and 36.2), issued by the American Psychiatric Association^{1,2} and in the *International Classification of Disease—Edition 10* criteria for research (ICD-10), issued by the World Health Organization³ (see Table 36.3). In these criteria, dementia is a syndrome characterized by a decline in memory and other intellectual functions (e.g. orientation, visuospatial abilities, language, thinking, executive function, problem solving, apraxia, agnosia). It is often accompanied by changes in behaviour or personality (e.g. loss of initiative, emotional lability, irritability, apathy, coarsening of social behaviour, change in mood). These latter symptoms are mandatory for a diagnosis in ICD-10, diagnostic in the presence of memory dysfunction in DSM-III-R and not included in DSM-IV.

An interesting difference between DSM-III-R and DSM-IV is that DSM-III-R requires impairment in short *and* long-term memory, while DSM-IV states that the impairment should include impairment in either short *or* long-term memory. In DSM-IV, in contrast to DSM-III-R and ICD-10, the criteria for dementia are integrated with the criteria for different types of dementia (such as Alzheimer’s disease and vascular dementia). It is thus not permitted to diagnose the dementia syndrome *per se*, but the subcriteria for dementia are identical for all types.

The symptoms of dementia are on a continuum with normal behaviour, which often makes it difficult to know where the line should be drawn between normal function and mild dementia. This dimensional rather than categorical character makes mild dementia difficult to separate from normal ageing⁴. Fairly small differences in criteria may have large effects on the prevalence rates. Mowry and Burvill⁵ found a variation in the prevalence of mild dementia ranging from 3% to 64% when different criteria were used on the same population. Different criteria also diagnosed different individuals. If a decline from a previously higher level can be shown (by obtaining information from key informants or by following the patients over time), the validity may be higher⁶. The DSM-III-R, DSM-IV and ICD-10 use the degree of social consequences of the disorder [“sufficient to interfere with everyday activities” (ICD-10) and “significant impairment in social or occupational functioning representing a significant decline from a previous level of functioning” (DSM-IV)] as the criterion for demarcating normal from abnormal behaviour.

The modern concept of dementia does not imply anything about prognosis, i.e. the course may be progressive, static, fluctuating or even reversible.

DIFFERENT TYPES OF DEMENTIA

A dementia syndrome may be caused by more than 70 diseases, the most common being Alzheimer’s disease and vascular dementias.

Alzheimer’s Disease

The neuropathology of Alzheimer’s disease (AD) is characterized by a marked degeneration of the neurons and their synapses and the presence of extensive amounts of extracellular senile plaques and intracellular neurofibrillary tangles in certain areas of the brain. The typical insidious onset and gradually progressive course is emphasized in the National Institute of Neurological and Communicative Disorders and Stroke and the Alzheimer’s Disease and Related Disorders Association (NINCDS–ADRDA) criteria (Table 36.4), in DSM-III-R and in DSM-IV but not in ICD-10. Memory disturbance is the most prominent early symptom, but slight impairment of visuospatial functioning, language and concentration may occur. In the later stages, the symptomatology is more widespread. The NINCDS–ADRDA criteria, DSM-III-R, DSM-IV and ICD-10 require that the diagnosis of AD should be made in the absence of diseases that, in and of themselves, could account for the progressive deficits in memory and cognition. Possible AD, according to the NINCDS–ADRDA criteria, may be diagnosed in the presence of other diseases if they are not judged to have caused the dementia.

Alzheimer’s disease is categorized into an early- and a late-onset form in DSM-IV and ICD-10, based on whether onset occurred before or after age 65 years. ICD-10 also specifies that the early-onset type should have a relatively rapid onset and progression or aphasia, agraphia, alexia, acalculia or apraxia, while the late-onset type should have a very slow, gradual onset and progression or predominance of memory impairment. Another subdivision is between familial AD (FAD) and sporadic AD. FAD has an autosomal dominant inheritance. Almost all cases with FAD have an early onset, while most cases of sporadic AD occur late in life. Familial clustering may occur also in sporadic AD⁸.

Table 36.1 Dementia (DSM-IV) (adapted)

-
- A1 Memory impairment (impaired ability to learn new information *or* to recall previously learned information)
and
- A2 One (or more) of the following cognitive disturbances:
(a) Aphasia
(b) Apraxia
(c) Agnosia
(d) Disturbance in executive functioning
- B The cognitive deficit in A1 and A2 *each* cause significant impairment in social *or* occupational functioning *and* represent a significant decline from a previous level of functioning
-

American Psychiatric Association².

Table 36.2 Dementia (DSM-III-R) (adapted)

-
- A Demonstrable evidence of impairment in short *and* long-term memory
and
- B At least one of the following:
(i) Impairment in abstract thinking
(ii) Impaired judgement
(iii) Other disturbances of higher cortical function, such as:
aphasia, apraxia, agnosia, constructional difficulty
(iv) Personality change
- C The disturbance in A + B significantly interferes with work *or* usual social activities *or* relationships with others
- D Not occurring exclusively during the course of delirium
- E Either (1) there is evidence from the history, physical examination, or laboratory tests of a specific organic factor (or factors) judged to be etiologically related to the disturbance, or (2) in the absence of such evidence, an etiologic organic factor can be presumed if the disturbance cannot be accounted for by any non-organic mental disorder
-

American Psychiatric Association¹.

Table 36.3 ICD-10 Criteria for dementia. Definition of dementia in the ICD-10 (adapted)

-
- G1 There is evidence of each of the following:
(1) A decline in memory (at least) sufficient to interfere with everyday activities, though not so severe as to be incompatible with independent living
(2) A decline in other cognitive abilities characterized by deterioration in judgement and thinking, such as planning and organizing, and in the general processing of information (at least) sufficient to cause impaired performance in daily living, but not to a degree that makes the individual dependent on others
- G2 Awareness of the environment (i.e. absence of clouding of consciousness)
- G3 There is a decline in emotional control or motivation, or a change in social behaviour manifest as at least one of emotional lability, irritability, apathy or coarsening of social behaviour
- G4 The symptoms in criterion G1 should have been present for at least 6 months
-

World Health Organization³.

Table 36.4 NINCDS-ADRDA Criteria for Alzheimer's disease

-
1. Probable Alzheimer's disease:
Dementia
Deficits in two or more areas of cognition
Progressive worsening of memory and other cognitive functions
No disturbance of consciousness
Onset between ages 40 and 90
Absence of systemic disorders or other diseases that in and of themselves could account for the progressive deficits in memory and cognition
2. Possible Alzheimer's disease:
Dementia
Variations in the onset, in the presentation, or in the clinical course
May be made in the presence of a second systemic or brain disorder sufficient to produce dementia, which is not considered to be the cause of the dementia
Should be used in research studies when a single, gradually progressive severe cognitive deficit is identified in the absence of other identifiable cause
-

McKhann *et al.*²⁹.

Vascular Dementia

Vascular dementia is a dementia caused by different forms of cerebrovascular disorder (CVD)^{9,10}, most often stroke and ischaemic white matter lesions (WMLs). The Hachinski Ischemic Score¹¹ was the most widely used instrument for the diagnosis of

vascular dementia, or rather multi-infarct dementia (MID), from the 1970s to the early 1990s. It consists of a symptom checklist that incorporates some of the symptoms that are believed to be essential in MID, such as abrupt onset, stepwise deterioration, fluctuating course, a history of stroke, and focal neurological symptoms and signs. The assumption was that MID was caused

by embolic phenomena, so that the onset of the clinical condition would be sudden and acute. Subsequent further emboli would produce other sudden deteriorations, perhaps followed by some improvement as areas of brain oedema resolved and some function was restored.

WMLs refer to the histopathological picture of diffuse demyelination with incomplete infarction in subcortical structures of both hemispheres, and arteriosclerotic changes with hyalinization or fibrosis of the small penetrating arteries and arterioles in the white matter⁹. These lesions may appear as low-density areas on computed tomography (CT) scans and as hyperdense areas on magnetic resonance imaging (MRI). The cognitive decline in subjects with WMLs has been suggested to be caused by a disconnection of subcortical–cortical pathways, producing a decline in abilities related to subcortical or frontal lobe structures.

Memory impairment is mandatory for the diagnosis of vascular dementia in ICD-10, DSM-III-R and DSM-IV. This is not ideal to describe the cognitive dysfunction in vascular dementia, where intellectual impairment may be substantial while memory dysfunction is mild¹². The ICD-10 requires that “deficits in higher cognitive functions are unevenly distributed” and DSM-III-R that there is “a patchy distribution of deficits (i.e. affecting some functions, but not others) early in the course”. The latter was, however, no longer included in the DSM-IV.

Although stroke increases the risk of developing dementia several-fold^{13,14} the contributions of a stroke or an infarct to the clinical symptoms of dementia are not always easy to elucidate. Stroke may be the main cause of dementia in an individual, it may be the event that finally overcomes the brain’s compensatory capacity in a subject whose brain is already compromised by Alzheimer pathology, albeit not yet clinically manifest, and in many instances minor manifestations of both disorders which individually would not be enough to produce dementia may produce it together¹⁵. Sometimes the presence of stroke in a patient with AD may be coincidental. Most criteria leave it to the clinician to make the decision whether the cerebrovascular disease “may be judged to be aetiologically related to the dementia” (ICD-10, DSM-III-R, DSM IV).

In most criteria the definition of CVD is based on history or findings of focal neurological upper motor neuron symptoms/signs, or brain imaging findings of CVD. DSM-IV (Table 36.5) gives examples of signs, while the ICD-10 (Table 36.6) specifically requires that at least one should be: (1) unilateral spastic weakness of the limbs; (2) unilateral increased tendon reflexes; (3) extensor plantar response; or (4) pseudobulbar palsy.

The DSM-IV specifies that there should be signs *and* symptoms *or* laboratory evidence indicative of CVD (e.g. multiple infarctions involving the cortex and underlying white matter) that are judged to be aetiologically related to the disturbance, while ICD-10 requires that there should be evidence from history,

Table 36.5 DSM-IV Vascular dementia

A/B	General criteria for dementia
C	Focal neurological signs and symptoms, e.g: Exaggeration of deep tendon reflexes Extensor plantar response Pseudobulbar palsy Gait abnormalities Weakness of an extremity <i>or</i> Laboratory evidence of cerebrovascular disease, e.g. Multiple infarctions involving cortex and underlying white matter that are judged to be etiologically related to the disturbance
D	Do not occur exclusively during delirium

American Psychiatric Association².

Table 36.6 ICD-10 Criteria for vascular dementia (adapted)

G1	The general criteria for dementia (G1–G4) must be met
G2	Deficits in higher cognitive function are unevenly distributed, with some findings affected and others relatively spared
G3	There is clinical evidence of focal brain damage, manifest as <i>at least</i> one of the following: (1) Unilateral spastic weakness of the limbs (2) Unilaterally increased tendon reflexes (3) An extensor plantar response (4) Pseudobulbar palsy
G4	There is evidence from the history, examination, <i>or</i> tests of a significant cerebrovascular disease, which may reasonably be judged to be etiologically related to the dementia

World Health Organization³.

examination *or* tests of a significant CVD, which may be reasonably judged to be aetiologically related to the dementia (e.g. history of stroke or evidence of cerebral infarction). In the National Institute of Neurological Disorders and Stroke and the Association Internationale pour la Recherche et l’Enseignement en Neurosciences (NINDS–AIREN) criteria¹⁰ (Table 36.7), a diagnosis of probably vascular dementia requires that focal signs consistent with stroke *and* relevant CVD by brain imaging should be present. Tatemichi, one of the authors of the NINDS–AIREN criteria, and his colleagues published a modified version¹⁶, in which this criterion was changed to focal signs consistent with stroke *or* relevant CVD by brain imaging. The first criterion is probably too strict and underestimates the occurrence of VaD; the latter criterion may be too broad. The NINDS–AIREN criteria recommend that a diagnosis of “possible” vascular dementia may be made in the presence of dementia with focal neurological signs in patients in whom brain imaging studies are missing; or in the absence of a clear temporal relationship between dementia and stroke; or in patients with subtle onset and variable course. This means that if CVD is present in a patient with dementia, VaD is likely to be diagnosed, which might overestimate the occurrence of this type of dementia. Furthermore, the interpretation of a single stroke leading to dementia probably differs between centres, and may be one reason for the disparate results regarding the prevalence of vascular dementia.

Table 36.7 The NINDS–AIREN criteria

Probable Vascular Dementia	
1.	Dementia
2.	Cerebrovascular disease (a) Focal signs consistent with stroke <i>and</i> (b) Relevant CVD by brain imaging: Multiple large-vessel infarcts Single strategically placed infarct Multiple lacunes (basal ganglia, white matter) Extensive periventricular white matter lesions
3.	Relationship between (1) and (2): (a) Dementia onset within 3 months following stroke (b) Abrupt deterioration in cognitive functions (c) Fluctuating stepwise progression
Possible Vascular Dementia	
1.	Dementia
2.	Cerebrovascular disease Focal signs consistent with stroke
3.	Absence of relationship between (1) and (2) (a) Dementia onset more than 3 months following stroke (b) Subtle onset or variable course

The temporal relationship between stroke and onset of dementia is often thought to strengthen the possibility that the two disorders are aetiologically related. The NINDS–AIREN criteria suggest an arbitrary limit of 3 months for onset of dementia after stroke. However, a stroke which occurred years before may still indicate the presence of CVD.

VAD may be underdiagnosed, as sometimes the onset is insidious, the course gradual, the infarctions clinically silent and the infarcts not detectable by CT of the brain^{17,18}. VAD may be overdiagnosed, as the presence of stroke, WMLs or other CVD does not necessarily mean that they are the cause of the dementia¹⁷. Often AD becomes a diagnosis by exclusion, and the diagnosis of VAD will be assigned if the patient has a history of CVD. This leads to a situation where the dementias will not infrequently be divided into one group with stroke and one without, giving negative associations between risk factors for stroke and AD, and positive associations with VAD¹⁷.

Even the histopathological diagnoses of AD and VaD are uncertain. Extensive histopathological signs of AD^{19,20} and VAD^{19,21} have been found in persons who show no clinical signs of dementia during life. A considerable proportion of subjects fulfilling the diagnosis of probable NINCDS–ADRDA criteria for AD or probable NINDS–AIREN for VAD have mixed pathologies^{22,23}. CVD may increase the possibility that individuals with AD lesions in their brains will express a dementia syndrome²⁴, but some workers have suggested that there may be a causal link between AD and VAD²⁵.

Frontotemporal Dementia

Frontotemporal dementia is a neurodegenerative disease characterized by neuronal loss in the frontal and temporal lobes. Criteria for frontotemporal dementia was proposed by the Lund and Manchester Groups in 1994²⁶, and revised in 1998²⁷. The latter describes five core diagnostic features that must be present: (a) insidious onset and gradual progression; (b) early decline in social interpersonal conduct; (c) early impairment in regulation of personal conduct (e.g. social disinhibition); (d) early emotional blunting (such as inertia and loss of volition); and (e) early loss of insight. It also includes supportive diagnostic features, which are not present in all patients: (a) behavioural disorder; (b) speech and language alterations (economical output, reduced number of words, asponaneity, press of speech, stereotypy, echolalia, mutism, perseveration); (c) physical signs (primitive reflexes, incontinence, akinesia, rigidity, tremor, low or labile blood pressure); (d) investigations showing impairment in frontal lobe tests in the absence of severe amnesia, aphasia or perceptual disorder, normal EEG, and frontal or anterior temporal abnormality on CBF. Other common symptoms are stereotyped behaviour and motor perseverations. Cognitive deficits occur mainly in attention, abstraction, planning and problem solving, while memory is relatively well-preserved in the early phase²⁷. In DSM-IV and ICD-10, this type of dementia is classified under the heading Dementia due to Pick's disease, while Pick's disease is a subtype of frontotemporal dementia in the criteria from Neary *et al*²⁷. According to the ICD-10, the general criteria for dementia should be met, onset should be slow with steady deterioration, memory and parietal lobe functions should be relatively preserved in the early stages, and at least two symptoms should be either emotional blunting, coarsening of social behaviour, disinhibition, apathy or restlessness, and aphasia. The problem with the emphasis on memory impairment in the general criteria for dementia in this disorder is evident.

Two other clinical syndromes of frontotemporal degeneration are progressive nonfluent aphasia and semantic dementia, which are disorders of language. Patients with Alzheimer's disease,

vascular dementia, and some other brain disorders may also exhibit symptoms of the frontal lobe type during the course of their disorders.

Lewy Body Dementia

Lewy body disease is a neurodegenerative dementia characterized by Lewy body formation in the brain stem and cerebral cortex. It has been reported that as much as 20% of demented patients coming to autopsy exhibit these changes. Criteria for Lewy body dementia are lacking in DSM-IV and ICD-10, but were proposed by McKeith *et al*²⁸ in 1992. These include: (a) fluctuating cognitive impairment affecting both memory and higher cortical functions; (b) at least one of (i) visual and/or auditory hallucinations, (ii) mild spontaneous extrapyramidal features (mainly rigidity and hypokinesia) or a neuroleptic sensitivity syndrome (i.e. exaggerated adverse response to standard doses of neuroleptic medications), or (iii) repeated unexplained falls and/or transient clouding or loss of consciousness; (c) despite the fluctuating pattern the clinical features persist over a long period of time; (d) exclusion of any underlying physical illness adequate to explain the fluctuating cognitive state, and of past history of stroke or ischaemic brain damage on brain imaging. The clinical presentation often also includes paranoid ideations and depression.

Subcortical Dementia

A special subtype of dementia is subcortical dementia. This type of dementia is seen in subcortical disorders, such as Parkinson's disease with dementia, Huntington's disease, supranuclear palsy, Lewy body disease and subcortical ischaemic WMLs. The dominating symptoms are psychomotor retardation, emotional bluntness, akinesia and slight memory disturbance, which may be helped by cues.

Secondary Dementias

Secondary dementias are caused by conditions with a known aetiology where dementia is generally not a core symptom, but may occur in some patients. Traditionally, vascular dementia is not classified among the secondary dementias, while subdural haematomas, normal pressure hydrocephalus, Creutzfeldt–Jakob disease, brain tumours, metabolic disorders and deficiency states are treated as secondary dementias.

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Cross-national Inter-rater Reliability of Dementia Diagnosis

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DSM-IV¹, ICD-10² and other glossaries have proved successful in promoting a common approach to psychiatric diagnosis. In field trials of DSM-III-R, for example, psychiatrists achieved concordance rates for diagnosing dementia of 0.91, where 1.0 represents complete agreement³.

It cannot be assumed, however, that similar performances will be achieved in “real world” practice. Most studies of diagnostic reliability involve assessments by experienced clinicians of cooperative, physically healthy old people who are either “normal” controls or “pure cases” of dementia. In reality, cognitive function lies on a spectrum and assessment is often complicated by limited education, deafness, poor vision, physical illness, anxiety or depression.

In a study of five research teams in Australia, Germany, The Netherlands, the UK and the USA, each centre contributed 20 written vignettes of elderly persons encountered in clinics or community surveys⁴. No exclusions were made because of

medical, neurological or psychiatric complications. The vignettes were brief and highly structured. The contents included subjects' demographic details, medical and psychiatric history, abbreviated cognitive test results and an informant's report of cognitive, personal and social performance.

When 13 researchers applied DSM-III-R criteria to the 100 vignettes, within-team levels of diagnostic agreement were high, ranging from kappa 0.72 in the centre with the least joint training to 0.86 in the centre with most. Between-team agreement was lower but still acceptable, with kappas ranging from 0.74 to 0.83. Some elements of DSM-III-R were easier to apply consistently than others. Mean percentage agreement was highest for social and occupational dysfunction (94%) and lowest for impairment of higher cortical function (87%). Concordance was also significantly higher for cognitively intact (98%) and severely impaired (96%) persons than for those with minimal (80%) and mild (82%) degrees of dementia.

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Cross-national Inter-rater Reliability of Dementia Diagnosis

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DSM-IV¹, ICD-10² and other glossaries have proved successful in promoting a common approach to psychiatric diagnosis. In field trials of DSM-III-R, for example, psychiatrists achieved concordance rates for diagnosing dementia of 0.91, where 1.0 represents complete agreement³.

It cannot be assumed, however, that similar performances will be achieved in “real world” practice. Most studies of diagnostic reliability involve assessments by experienced clinicians of cooperative, physically healthy old people who are either “normal” controls or “pure cases” of dementia. In reality, cognitive function lies on a spectrum and assessment is often complicated by limited education, deafness, poor vision, physical illness, anxiety or depression.

In a study of five research teams in Australia, Germany, The Netherlands, the UK and the USA, each centre contributed 20 written vignettes of elderly persons encountered in clinics or community surveys⁴. No exclusions were made because of

medical, neurological or psychiatric complications. The vignettes were brief and highly structured. The contents included subjects' demographic details, medical and psychiatric history, abbreviated cognitive test results and an informant's report of cognitive, personal and social performance.

When 13 researchers applied DSM-III-R criteria to the 100 vignettes, within-team levels of diagnostic agreement were high, ranging from kappa 0.72 in the centre with the least joint training to 0.86 in the centre with most. Between-team agreement was lower but still acceptable, with kappas ranging from 0.74 to 0.83. Some elements of DSM-III-R were easier to apply consistently than others. Mean percentage agreement was highest for social and occupational dysfunction (94%) and lowest for impairment of higher cortical function (87%). Concordance was also significantly higher for cognitively intact (98%) and severely impaired (96%) persons than for those with minimal (80%) and mild (82%) degrees of dementia.

Agreement was higher for “yes–no” DSM-III-R diagnoses than for the multilevel Clinical Dementia Rating (CDR) scale, in which six domains (memory, orientation, judgement, community affairs, home and hobbies, and personal care) are each rated on a five-point scale⁵. Kappa levels ranged from 0.61 to 0.76 within teams and from 0.50 to 0.69 between teams. Personal care was rated most consistently (85% mean agreement) and community affairs least so (74%). As with DSM-III-R, concordance was higher for cognitively intact (95%) and severely impaired (84%) persons than for those with minimal (78%) and mild (68%) dementias.

Univariate analyses of CDR ratings pointed to lower agreement levels for persons described as physically ill, deaf, partially sighted, anxious or depressed. However, multivariate analysis detected only two main effects: dementia severity and physical illness. Other variables made no significant independent contribution.

These findings suggest that dementia can be diagnosed with acceptable reliability in community surveys. Agreement is likely to be higher, though, when teams train intensively and use instruments that require simple “yes–no” choices. The reduction in agreement associated with physical illness is important given

the high co-morbidity of physical and mental illness in representative elderly populations.

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Early Detection

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IS EARLY DETECTION IMPORTANT?

Why try to detect dementia early in its course? Since the advent of technologies for the early detection of disease, it has become important to ask whether early recognition is worthwhile, and for whom. Can it be shown that a lower level of morbidity is achieved in a population that has been screened, compared to others that were not screened? Is the quality of life of cases improved by their early diagnosis; or could early detection of dementia “seriously damage your health”, as has been found for other disorders¹? For a start, evaluation of the early detection of dementia would have to take note of the 10 principles of screening, as listed by Wilson and Jungner². These include the requirement that there be an accepted form of treatment for persons once detected, that facilities for diagnosis and treatment be available to the population being screened, and that a suitable test be available for detecting the disease in its early stages. Clearly, screening for early dementia is a procedure where none of these requirements has yet been met.

Cooper and Bickel³ have nevertheless indicated some of the advantages that screening or early detection could bring. They point out, first, that the biggest gap between those receiving specialist psychiatric care and the total volume of morbidity in a general population is amongst its elderly; and that mental disorders in this age group are probably under-recognized by general physicians. Second, a proportion of cases detected as dementia have reversible conditions such as depressive disorder, normal pressure hydrocephalus, metabolic disorders or brain tumours. Third, early detection can be a preliminary not to curative treatment but “to intervention aimed at reducing disability and postponing the need for institutional care”. Such intervention is well placed with the general practitioner and the person’s family. It can then be added that, for the general practitioner, the advantages of early detection are appreciable: the possible causes of the cognitive or behavioural deterioration may need to be pursued. Where co-morbidity emerges, as it commonly will, the physician’s awareness that dementia is present will prove useful in assessment and continuing management; and the presence of dementia may influence the choice of medication.

There is one further reason for early detection, although this is not to the individual patient’s immediate benefit. For research on dementia, it is of great importance to know about the earliest symptoms and signs before these become buried by the dementia itself. Without this information, and without knowing the clinical course of mild cases, it may not be possible to improve the specificity of screening, and to distinguish between mild cognitive

decline and normal ageing⁴. For population-based research, where some early cases will inevitably be identified, Brayne *et al.*⁵ recommend that a two-stage design should ideally be used, along with a third assessment that serves as a gold standard: evidence of progression of the dementia; or neuropathology at post mortem.

HOW IS EARLY DETECTION ACHIEVED?

Early detection will usually mean the dementia is of only mild severity. It is an advantage, therefore, that the diagnostic criteria for mild dementia have been specified in both ICD-10⁶ and DSM-IV⁷. In ICD-10, the declining memory and information processing causes impaired performance in daily living, but not to a degree that is incompatible with independent living. Explicit criteria are given for the diagnosis of mild dementia.

Early detection can be carried out at three levels: in the community, in primary care settings, and in hospitals. In the community, screening is conducted only as part of research studies, and has not yet been used in a way similar to other routine screening for disease. In primary care, early detection is at present conducted informally and is based largely on the initiative and clinical skill of the practitioner. It is not clear what most commonly prompts the physician into considering a diagnosis of early dementia. Rarely it would be, say, all persons aged 70 years and over consulting in a given period, but rather those who prompt the physician’s concern. Commonly, it is the patient’s family who have first detected a deterioration in cognition or behaviour. The present consensus is that, in the absence of some indication, efforts to detect early dementia in general practice are unwarranted. But there may be a place for routinely obtaining a base-line measure of cognitive performance against which subsequent assessments can be placed. There is as yet no place for annual repetition of the assessment.

Tests for Early Detection of Dementia

In hospitals and clinics, the realities of clinical practice are that early detection of dementia is achieved in one of two ways. In the more common style, the clinician obtains a history from others that a *decline* in cognitive performance and/or behaviour has taken place. To this is added some non-systematic cognitive assessment of the patient, leading to a conclusion on whether or not early dementia is present. Clearly, other clinical features,

such as cerebrovascular disease, would commonly play a part in this.

In the second situation, the assessment is formal and partly quantitative. A large number of standardized clinical instruments and neuropsychological tests for this are now available. They are described in Section DIII of this volume. For the purpose of *early* detection, two points need to be emphasized. Firstly, some of the tests are open to educational or cultural bias, so that they can generate false-positive results in some sociodemographic contexts, possibly to the patient's detriment. Secondly, in this writer's opinion, many tests are focused on cognitive function, ignoring changes in behaviour—yet the latter are an important clue to early dementia. The Psychogeriatric Assessment Scales developed by Jorm *et al.*⁸ have gone some way to redress this imbalance (*see* Chapter 27).

Limitations

To detect dementia early in its course, and to do so with a high level of accuracy in different social and educational groups, is not a straightforward task. The only means currently available are those clinical instruments described in Section DIII. There is no biomedical test with both portability and greater accuracy than these instruments. The clinical instruments all have a number of limitations. First, the mental status questionnaires are brief, and can act only as screening instruments that sort individuals into different levels of probability of being a case of dementia. A questionnaire cannot be expected to do more than this. Second, the instrument must be acceptable; yet elderly people may dislike extensive cognitive testing, particularly if it shows them up as defective in performance. Third, whether it is a brief questionnaire or a clinical examination, the reliability and validity have to be high. The latter means achieving good levels of sensitivity and specificity. Fourth, and closely related to validity, there is the problem of bias against low intelligence or poor education. It is highly likely that this causes some false-positives to appear in the course of screening.

Whatever the method used for early detection, there are three further issues to consider. Not all the cases detected will progress^{9,10} and it is hard to predict to whom this will happen.

ORGANIC DISORDERS

Rosenman¹¹ found very poor predictive validity for five well-established methods for making this diagnosis. Cooper and Bickel³ argue that there are no good grounds for asking elderly persons to subject themselves to extensive investigations when some will undergo no further deterioration. A further impediment is the cost and service burden from investigating all the possible cases of dementia generated from a national screening programme. Eastwood and Corbin¹² have argued that the cost would be prohibitive. This is likely to be the case even for the older section of the community, where the frequency of secondary dementias is known to be lower than in younger adults. Lastly, there may be unexpected adverse consequences of screening. The belief may be false that early detection can contribute to prevention. O'Connor *et al.*¹³ found, unexpectedly, that screening for early or mild dementia increased the likelihood of entry into residential care.

WHAT POSSIBLE SOLUTIONS ARE THERE?

Routine screening for dementia at the community level cannot yet be defended. What is required instead is work to evaluate

its impact in the manner advocated by Cooper and Bickel³. These authors have emphasized the need for research into the feasibility and effectiveness of early detection as a first step towards preventive action. They also argue that programmes for early detection will be successful only if they are incorporated into the work of general practitioners, community nurses and other health professionals. It is there, and not the total community of elderly, where early detection needs first to be attempted and evaluated to see what benefits, if any, it brings.

At a technical level, it has to be accepted that early detection of cognitive decline or dementia by brief screening methods has at least two unavoidable imperfections: first, there will be some error, whereby the screening test misclassifies a proportion of individuals; second, some of this error will be attributable to low intelligence in the respondent, or educational bias in the test. Since both of these are likely to produce false-positives rather than negatives, the problem can usually be overcome by more detailed clinical assessment and history in a two-phase design. In research settings, another desirable strategy in early detection is to have a second assessment after an appropriate lapse of time. This is the most certain way to ensure that deterioration has indeed taken place, and that it has progressed. Early detection of dementia is currently dependent on clinical information and on cognitive performance related to daily life. Assessment of this is now remarkably satisfactory, although the validity of the main clinical instruments has yet to be demonstrated in community instead of hospital samples. Because dementia has very explicit clinical manifestations and associated impairments, its early detection is likely to be by clinical means for some time to come. For the present, early detection brings no benefit to the elderly in the general population. But the situation may soon change in the face of current advances in the molecular biology of Alzheimer's disease, where it is conceivable that early detection may become justified for genetically high-risk individuals for whom a pharmacological intervention could be beneficial¹⁴. It is from these advances that far-reaching consequences for clinical practice are now imminent.

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Dementia Epidemiology: Prevalence and Incidence

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Prevalence is the proportion of cases of a disease present in a population at any one time, while incidence is the rate of occurrence of new cases over a given period of time, usually 1 year. Prevalence is a function of both the incidence of disease and its duration: the prevalence of a disease will rise if the rate of new cases increases or if the average case survives longer. Prevalence is useful for assessing the likely need for service provision. However, for purely scientific purposes, such as assessing risk factors, incidence is preferred over prevalence. The reason is that any differences between groups in prevalence may be due to differences in either incidence or disease duration.

The notions of prevalence and incidence are based on the assumption that a population can be neatly divided into cases and non-cases. However, for dementia this division is not straightforward. There is a gradation from normal cognitive ageing through to severe dementia, without clear breaks to define where normality ends and dementia begins. The threshold for dementia is usually defined in terms of interference with daily living, but even this is a fuzzy boundary. Furthermore, prevalence and incidence studies typically examine different levels of severity, described as “mild”, “moderate” or “severe”, but these descriptors are not always used consistently to divide up the continuum of severity. Various diagnostic criteria for dementia are known to divide the continuum in different ways, which can result in very different prevalence and incidence rates. For example, Erkinjuntti *et al.*¹ examined the rates of dementia in the same sample using six different sets of diagnostic criteria. They found that the percentage with dementia varied from 3.1% using the ICD-10 criteria to 29.1% using DSM-III. Thus, there are no “true” prevalence or incidence rates for dementia, but rather various rates dependent on the definition of dementia used.

PREVALENCE OF DEMENTIA

The number of prevalence studies is now very large and several meta-analyses have been carried out to pool the data for those that give rates for specific age groups (e.g. 65–69 years). The first meta-analysis, by Jorm *et al.*², involved fitting a statistical model to data from 22 studies published between 1945 and 1985. They found that methodological differences between studies contributed to variation in prevalence rates. For example, studies that used a broad definition of dementia (to include all cognitive impairment) had rates 64% higher than those using a more narrow definition. They fitted an exponential statistical model to the data². The essence of this model is that prevalence rises

exponentially with age, doubling every 5.1 years, but the actual rates differ from study to study. Although there were differences between studies, it is possible to derive average rates across studies. These are shown in column 1 of Table 38.1. The exponential model is an adequate description up to age 90 but should not be applied above that age. If prevalence continued to double every 5.1 years above age 90, it would soon be greater than 100%, which is impossible. This limitation of the exponential model has led some researchers to use the logistic model, in which prevalence at first rises steeply, but then levels out to a maximum of 100%³. For prevalence rates of 0–50% the exponential and logistic curves are difficult to distinguish, and the exponential curve may be preferred because of its simplicity.

The second meta-analysis involved a pooling of data from 12 European studies dating 1980–1990, which used DSM-III or equivalent criteria⁴. This meta-analysis did not involve fitting a statistical model to the data or testing for the effects of methodological differences. Rather, the researchers simply divided the data from each study into 5 year age groups and pooled them. The results are shown in column 2 of Table 38.1. Despite the differences in approach, the results are remarkably close to those of Jorm *et al.*².

A third meta-analysis was carried out by Ritchie and Kildea⁵ (this superseded an earlier meta-analysis by Ritchie *et al.*⁶, which will not be described here). They were particularly interested in what happens to prevalence in extreme old age, in particular whether dementia is inevitable if a person lives long enough. Ritchie and Kildea⁵ pooled data from nine studies that included samples of elderly people aged over 80. These data are shown in column 3 of Table 38.1 and are very similar to the earlier meta-analyses up to age 90. Ritchie and Kildea⁵ fitted various curves to

Table 38.1 Prevalence rates (%) for dementia from three meta-analyses

Age group	Meta-analysis		
	Jorm <i>et al.</i> ²	Hofman <i>et al.</i> ⁴	Ritchie and Kildea ⁵
60–64	0.7	1.0	—
65–69	1.4	1.4	1.5
70–74	2.8	4.1	3.5
75–79	5.6	5.7	6.8
80–84	11.1	13.0	13.6
85–89	23.6*	24.5*	22.3
90–94	—	—	33.0
95–99	—	—	44.8

* Rates for ages 85+.

the data, including exponential and logistic models. The best fit was by a modified logistic curve in which prevalence levelled off at around 40% at age 95. The authors concluded that dementia is not inevitable in extreme old age. However, this conclusion has been criticized by McGee and Brayne⁷ because it was based on prevalence rather than incidence data. A decrease in survival with dementia in very old age could explain the flattening of the age-curve that was observed.

PREVALENCE OF ALZHEIMER'S DISEASE AND VASCULAR DEMENTIA

The clinical diagnosis of Alzheimer's disease and vascular dementia in community surveys involves additional problems beyond those in diagnosing global dementia, so such studies have been fewer. Nevertheless, several meta-analyses have attempted to integrate data on the issue.

The original meta-analysis of Jorm *et al.*² also examined seven studies that gave age-specific data on Alzheimer's disease and vascular dementia. One study could not be fitted well by the

exponential model, but the remaining six could. The prevalence of Alzheimer's was found to double every 4.5 years of age, while vascular dementia doubled every 5.3 years. In other words, the rise with age was steeper for Alzheimer's disease.

The pooling of data from European studies also examined specific dementing diseases⁸. The pooled prevalence rates for Alzheimer's disease from six studies were: 30–59 years, 0.02%; 60–69 years, 0.3%; 70–79 years, 3.2%; and 80–89 years, 10.8%. It was not possible to arrive at pooled rates for vascular dementia because of the variation across studies.

Later, Corrada *et al.*⁹ fitted a logistic model to data from 15 studies giving age-specific data on prevalence of Alzheimer's disease. They found considerable variation in rates between studies depending on the methodology used. However, the underlying trend was for an 18% increase in the odds for Alzheimer's disease with every year of age.

The most recent meta-analysis was carried out by the US General Accounting Office¹⁰ in response to controversy in that country about the number of people with Alzheimer's disease. A logistic curve was fitted to data from 18 studies with predominantly White populations. The data were grouped by sex and severity level of the dementia and are shown in Table 38.2. It can be seen that the results vary, depending on severity, and that females have a higher prevalence than males.

Table 38.2 Prevalence rates (%) for Alzheimer's disease according to a meta-analysis by the US General Accounting Office¹⁰

Age	All severity levels		Moderate–severe cases	
	Men	Women	Men	Women
65	0.6	0.8	0.3	0.6
70	1.3	1.7	0.6	1.1
75	2.7	3.5	1.1	2.3
80	5.6	7.1	2.3	4.4
85	11.1	13.8	4.4	8.6
90	20.8	25.2	8.5	15.8
95	35.6	41.5	15.8	27.4

INCIDENCE STUDIES

Incidence studies are much rarer than prevalence studies because they require longitudinal data and large sample sizes to arrive at age-specific rates. It is only fairly recently that sufficient studies have become available to permit meta-analysis.

Two meta-analyses have been published at around the same time. The first of these, by Jorm and Jolley¹¹, used data from 23 published studies. The incidence of both dementia and Alzheimer's disease was found to increase exponentially with age up to

Table 38.3. Incidence rates (%) for dementia from meta-analyses by Jorm and Jolley¹¹ and Gao *et al.*¹²

Age group	Jorm and Jolley ¹¹				Gao <i>et al.</i> ¹² 12 studies
	Europe (mild +)	Europe (moderate +)	USA (moderate +)	East Asia (mild +)	
60–64	—	—	—	—	0.11
65–69	0.91	0.36	0.24	0.35	0.33
70–74	1.76	0.64	0.50	0.71	0.84
75–79	3.33	1.17	1.05	1.47	1.82
80–84	5.99	2.15	1.77	3.26	3.36
85–89	10.41	3.77	2.75	7.21	5.33
90–94	17.98	6.61	—	6.61	7.29
95+	—	—	—	—	8.68

Mild+ results from USA and Moderate+ results from East Asia are missing from the table because insufficient data were found in the literature.

Table 38.4 Incidence rates (%) for Alzheimer's disease from meta-analyses by Jorm and Jolley¹¹ and Gao *et al.*¹²

Age group	Jorm and Jolley ¹¹					Gao <i>et al.</i> 8 studies
	Europe (mild +)	Europe (moderate +)	USA (mild +)	USA (moderate +)	East Asia (mild +)	
60–64	—	—	—	—	—	0.06
65–69	0.25	0.10	0.61	0.16	0.07	0.19
70–74	0.52	0.22	1.11	0.35	0.21	0.51
75–79	1.07	0.48	2.01	0.78	0.58	1.17
80–84	2.21	1.06	3.84	1.48	1.49	2.31
85–89	4.61	2.26	7.45	2.60	3.97	3.86
90–94	9.66	4.77	-	-	-	5.49
95+	-	-	-	-	-	6.68

Moderate+ results from East Asia are missing from the table because insufficient data were found in the literature.

90 years, with no sign of levelling off. The incidence of vascular dementia showed similar trends, but the actual rates varied greatly from study to study. There was no sex difference in dementia, but women tended to have a higher incidence of Alzheimer's disease in very old age, and men a higher incidence of vascular dementia at younger ages. There were also regional differences, with East Asian countries having a significantly lower incidence of dementia than Europe, and also tending to have a lower incidence of Alzheimer's disease. Tables 38.3 and 38.4 summarize the results for different regions and levels of severity.

The second meta-analysis, by Gao *et al.*¹², involved only the subset of 12 studies that used DSM-III criteria for dementia and NINCDS-ADRDA criteria for Alzheimer's disease. The data were fitted with a logistic model and a levelling of the rate of increase with age was found. Women were found to have a higher incidence of Alzheimer's disease than men. The estimated incidence rates are also shown in Tables 38.3 and 38.4. It can be seen that the rates of Gao *et al.*¹² are different from those of Jorm and Jolley¹¹ and the two meta-analyses came to different conclusions about whether there is a levelling in the rise with age. The difference arises because Gao *et al.*¹² pooled data from different regions and different levels of severity. Their rates fall in between those of Jorm and Jolley¹¹ for Mild+ and Moderate+ dementia. Deviations from an exponential rise can result if the studies contributing cases at the upper ages are examining milder dementia or are from regions with a lower incidence.

CONCLUSIONS

The prevalence and incidence of dementia rise exponentially with age up to age 90. There is no consensus about what happens at extreme ages, because of the limited data available, but some levelling in the rise is a possibility. Women probably have a higher prevalence and incidence of Alzheimer's disease. Conversely, men may be at greater risk of vascular dementia. There appear to be important regional differences, although the proper investigation

of these requires studies with identical methodologies in the various sites.

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Case-control Studies

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The case-control study is aimed at aetiology. Schlesselman¹ says that it has two distinctive features: it proceeds backwards from effect to cause by trying to identify exposures or other factors that led to a given disorder; and it uses a control or comparison group without the disorder, so that a causal effect for a given exposure can be supported or refuted. The first of these features is really what a clinician does in daily practice when taking a history, but as a rule, the clinician does not go on to determine how many *normal* persons have had the same exposure. The strength of the case-control method lies in these two features. It is on the basis of them that the fundamental comparison in the case-control study lies: the frequency of an exposure in the cases, and the frequency in the controls. Such a comparison is disarmingly simple, mainly because of the biases that can bring about misleading results. Some readable accounts can be found in Cole², Lilienfeld and Lilienfeld³, Feinstein⁴ and Anthony⁵. A non-technical overview has been set out by Henderson⁶. An entire issue of *Epidemiologic Reviews* devoted to the case-control method has provided an excellent conspectus of this powerful tool, including the diverse applications now being made of case-control designs for problem-solving in the health field, including evaluation of service interventions⁷.

In case-control parlance, “exposure” refers not only to environmental exposures, but to other properties of the individual, such as a family history of a particular disease or some other personal attribute. To identify an exposure that may contribute to the onset of a disorder, the investigator has firstly to choose a number of candidate exposures. This may be based on theory, on speculation, or on a mindless search. The first of these is particularly desirable, because it means that from the start there is some plausible biological or psychosocial basis for the putative effect. Speculation can be the vehicle for a gifted insight. The atheoretical examination of a large array of factors is undesirable and carries the risk of capitalization on chance.

CONDUCTING A CASE-CONTROL STUDY

There are five issues that deserve close attention:

1. The cases should be newly diagnosed, not ones which have been known for some time; and they should be representative of all incident cases in the population. If longer-established cases were used, the findings might be related more to factors influencing survival or chronicity than to aetiology (see below).
2. The cases should include no errors in diagnosis, which would lead to misclassification and therefore errors in estimating the relative risk for exposures.
3. The controls should either be matched demographically or be similar in overall attributes. Much thought needs to be accorded to the source of the controls if misleading biases are to be avoided.
4. In obtaining information on exposures from cases and controls, as well as from their informants, it is likely that selection effects will operate, causing information bias. That is, people may selectively recall certain experiences, or selectively report what they do recall. A likely example is a history of past head injury in Alzheimer’s disease, or any other situation where “effort after meaning” may operate. Ideally, the interviewers should themselves be blind to the purpose of the study, lest they unwittingly influence the information that they elicit.
5. A case-control study that has too few cases to provide a satisfactory estimate of risk is of little value. The sample size needed can be determined beforehand by establishing the minimum size of the effect to be demonstrated, and the frequency of exposure in the controls. The more the frequency of the exposure departs from 50% of the subjects, the more cases will be needed for an association to be shown.

The assessment of risk for an exposure is obtained by calculating its odds ratio as an approximation of the relative risk, and the 95% confidence intervals for that estimate (Schlesselman¹, p. 32 *et seq.*) (Henderson⁶, p. 16 *et seq.*). An odds ratio of 1.0 means that the exposure occurs as often in cases as in controls. The confidence interval should keep the estimate above unity if the exposure is to be accorded attention. It is misleading to report odds ratios without also giving their confidence intervals. For example, an odds ratio of 1.7 with a 95% confidence interval of 0.9–2.5 should be seen as a negative finding, because the lower limit is below unity.

RESULTS OF CASE-CONTROL STUDIES OF DEMENTIA

Nearly all the case-control studies of dementia have been focused on Alzheimer’s disease (AD). Within this diagnostic group, the studies have covered several categories, although not always making this explicit. The cases have often been heterogeneous in age of onset and, indeed, in age *since* onset. The latter introduces the problem of Neyman’s bias⁵ and factors related to survival after the onset of the dementia. In looking critically at case-control studies of dementia, their strengths and deficiencies can be seen by using the above points as a checklist, to assess the value that can be attached to each observation. Most studies have had only modest sample sizes. In case-control studies of AD, the

scientific significance of a result should rest on its replication by other workers on other samples; and on the development of biological evidence to support it as a risk factor in AD.

Risk and Protective Factors for Alzheimer's Disease

A comprehensive review of the evidence for a wide range of proposed risk factors, but also protective factors for AD, have been given by Jorm⁸ and Henderson and Jorm⁹.

Summarized, these are as follows:

Definite

- Age.
- Family history.
- Specific genetic mutations (for familial cases only).
- Apolipoprotein E ϵ 4 allele on chromosome 19.
- Down's syndrome.

Awaiting confirmation

- Regional or ethnic differences.
- α -2 Macroglobulin gene.
- Head injury (interaction with apoE?).
- Previous depressive disorder.
- Herpes simplex virus.
- Cerebrovascular disease.

Unlikely

- Aluminium in drinking water.
- Diet.
- Industrial solvents.
- Life stress.
- Electromagnetic fields.

Possible protective factors

- Education, high premorbid intelligence or both.
- Anti-inflammatory drugs (NSAIDs).
- Oestrogen.
- Smoking.
- Moderate wine drinking.

For vascular dementia, the review by Skoog¹⁰ lists the same risk factors as for stroke, namely hypertension, diabetes mellitus, advanced age, being male, smoking and cardiac disease. A

number of recent case-control studies have suggested that vascular factors may have a part to play in AD. There are as yet no case-control studies of Lewy body or other less common dementias, although there is no impediment other than ensuring accuracy of ascertainment and recruitment of sufficient cases. The same applies to other under-researched areas of geriatric psychiatry, such as the psychoses of late onset, where case-control studies could throw much-needed light on pathogenesis.

It does seem justifiable to continue to search for environmental exposures and other risk factors for AD, for vascular dementia and possibly the rarer dementias. Case-control research may in this way contribute not only to understanding their aetiology, but to finding preventive measures—the ultimate purpose of epidemiology. With the great expansion of the world's elderly, the social significance of this would be inestimable.

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Results from EURODEM Collaboration on the Incidence of Dementia

LJ Launer¹, for the EURODEM Incidence Research Group*

¹Erasmus University Medical Centre, Rotterdam, The Netherlands, and National Institutes of Health, Bethesda, MD, USA

In 1988, European investigators formed the EURODEM network to harmonize the protocols used in newly initiated population-based prospective studies on incident dementing diseases¹.

*Participants of the EURODEM Incidence Research Group: Department of Epidemiology and Biostatistics, Erasmus Medical School, The Netherlands (A. Hofman MD, L.J. Launer PhD, A. Ott MD, T. Stijnen PhD); Epidemiology, Demography, Biometry Program, National Institute on Aging, US (L.J. Launer PhD); Department of Psychiatry, Odense University, Denmark (K. Andersen MD, P. Kragh-Sorensen MD); Department of Psychiatry, Royal Liverpool University Hospital, UK (J.R.M. Copeland MD, M.E. Dewey PhD), INSERM Unit 330, France (J.F. Dartigues MD, L. Letenneur PhD); National Research Council Targeted Program on Ageing, Italy (L.A. Amaducci MD: now deceased); Institute of Public Health, Cambridge University, UK (C. Brayne MD); Department of Psychiatry, Zaragoza University (A. Lobo MD) and Department of Neurology, University of Navarra, Spain (J.M. Martinez-Lage MD).

Incident studies succeeded the case-control studies based on prevalent cases that were conducted in the 1980s². Studies based on incident cases are preferred to those based on prevalent cases, as the latter have several biases that affect the validity of their results³. Here we summarize the findings from the pooled EURODEM analyses on the frequency and risk for dementing disease in the elderly.

STUDY DESIGN

The pooled analyses are based on studies from Denmark⁴, France⁵, The Netherlands⁶ and the UK⁷. The sample includes

scientific significance of a result should rest on its replication by other workers on other samples; and on the development of biological evidence to support it as a risk factor in AD.

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Incident studies succeeded the case-control studies based on prevalent cases that were conducted in the 1980s². Studies based on incident cases are preferred to those based on prevalent cases, as the latter have several biases that affect the validity of their results³. Here we summarize the findings from the pooled EURODEM analyses on the frequency and risk for dementing disease in the elderly.

STUDY DESIGN

The pooled analyses are based on studies from Denmark⁴, France⁵, The Netherlands⁶ and the UK⁷. The sample includes

528 incident dementia cases and 28 768 person-years of follow-up. As a common core, each study included a population-based cohort of persons aged 65 years and older living in the community and nursing homes. Samples were drawn from defined geographic areas and either include all eligible individuals or individuals randomly selected within predefined strata. All studies contributed to the pooled analyses baseline data and one follow-up panel conducted after a fixed interval of about 3 years. The cohorts exclude the prevalent cases identified at baseline.

Dementia cases were identified in a two-stage procedure, whereby the total cohort was screened and screen-positive subjects underwent a detailed diagnostic assessment that included a clinical exam, neuropsychological testing and an informant interview. Dementia and vascular dementia were diagnosed according to DSM-III-R criteria⁸, Alzheimer's disease (AD) was diagnosed according to NINCDS-ADRDA criteria⁹.

RESULTS

AD comprised approximately 70% of all cases. Incident rates for dementia and AD were similar across studies. There was an increase with age in the incidence of all dementia and in particular AD. At 90 years of age and older the incidence of AD was 63.5 (95% CI, 49.7–81.0) per 1000 person-years. However, compared to men, women had significantly higher rates of AD after age 85 years. At 90 years of age, the rate of AD per 1000 person-years was 81.7 (63.8–104.7) in women and 24.0 (10.3–55.6) in men (Figure 1a,b)¹⁰. This translated into a cumulative risk for 65 year-old women to develop AD at the age of 95 years of 0.22 compared to 0.09 for men. These sex differences were not found in vascular

dementia: at 90 years of age, the incidence of vascular dementia was 15.9 (6.6–38.5) and 9.2 (4.3–19.6) per 1000 person-years in men and women, respectively.

In addition to sex and age, we investigated the association of AD to four risk factors that had been previously investigated in case-control studies^{1,2}. These risk factors were ascertained in a similar manner across studies. We found that low education increased the risk for dementia, specifically AD. However, the increased risk was detectable only in women: compared to those with high education, those with low education had a 4.3 (95% CI, 1.5–11.9) times, and those with middle education had a 2.6 (95% CI, 1.0–7.1) times increased risk for AD. There was no association of educational level to dementia among men¹¹. Contrary to a previous EURODEM analysis based on case-control studies², we did not find an increased risk for AD associated with head trauma. Previous reports based on prevalent cases suggested an inverse association of smoking with the risk for AD². In these current analyses based on incident cases, we found current smoking significantly *increased* the risk of AD. The risk associated with smoking was higher in men than women¹. Finally, we found that the association of AD to family history in two or more family members was weaker (OR 1.59 95% CI, 0.78–3.26) than previously estimated on the basis of case-control studies².

SUMMARY

Because we had a large sample, we were able to investigate whether gender modified the risk for AD, the most common form of late-life dementia. We found that women not only had a higher risk for AD than men, but that the relations of education and smoking to the risk for AD were different in men and women. These findings suggest that the risk for AD is altered either by sex-related biological or behavioral factors or by differences in cumulative survival.

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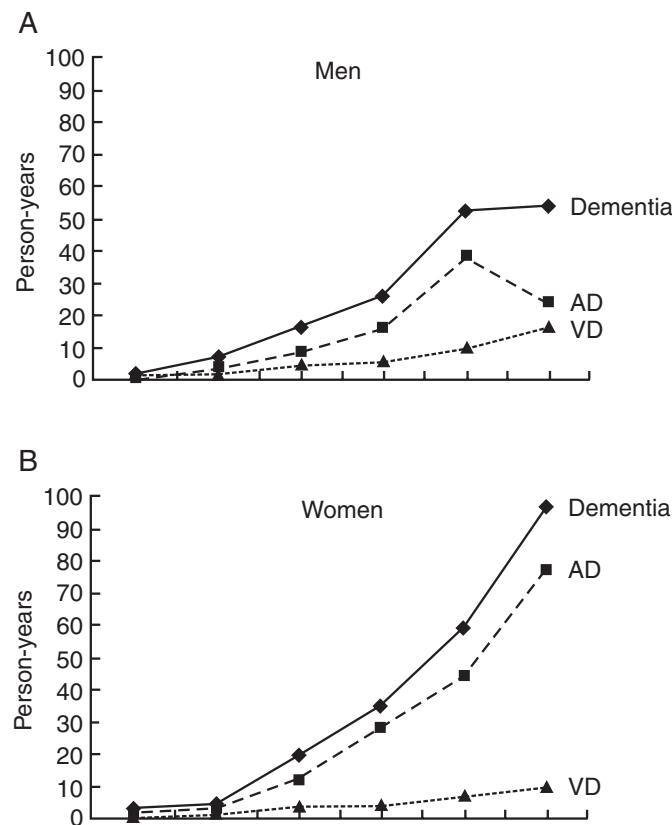


Figure 1 Incidence of dementia and major sub-types, Alzheimer's disease (AD) and vascular dementia (VD): EURODEM Studies. (A) men; (B) women

MRC/DoH Cognitive Function and Ageing Study

J. Nickson, C. F. M. McCracken and C. Brayne, on behalf of MRC CFAS

University of Cambridge, Cambridge, UK

The MRC Cognitive Function and Ageing Study (MRC CFAS) is a multi-centre prospective cohort study, set up in 1989 and funded by the MRC and Department of Health (DoH).

AIMS

CFAS aims to estimate the prevalence and incidence of cognitive decline and dementia and geographical variation; to determine the natural history of dementia, in particular the rate of progression of cognitive decline, including the distribution of the interval between identification of cognitive impairment and death; to identify factors associated with differing rates of

cognitive decline and with the risk of dementia; to determine the contribution of different underlying pathologies to rates of dementia, geographical variation and burden of disability; to evaluate the degree of disability associated with cognitive decline and service needs generated; to set up a brain and blood resource; and to provide a framework to support sub-studies.

METHODOLOGY

The study has five methodologically identical centres (Cambridge, Gwynedd, Newcastle, Nottingham and Oxford) and one

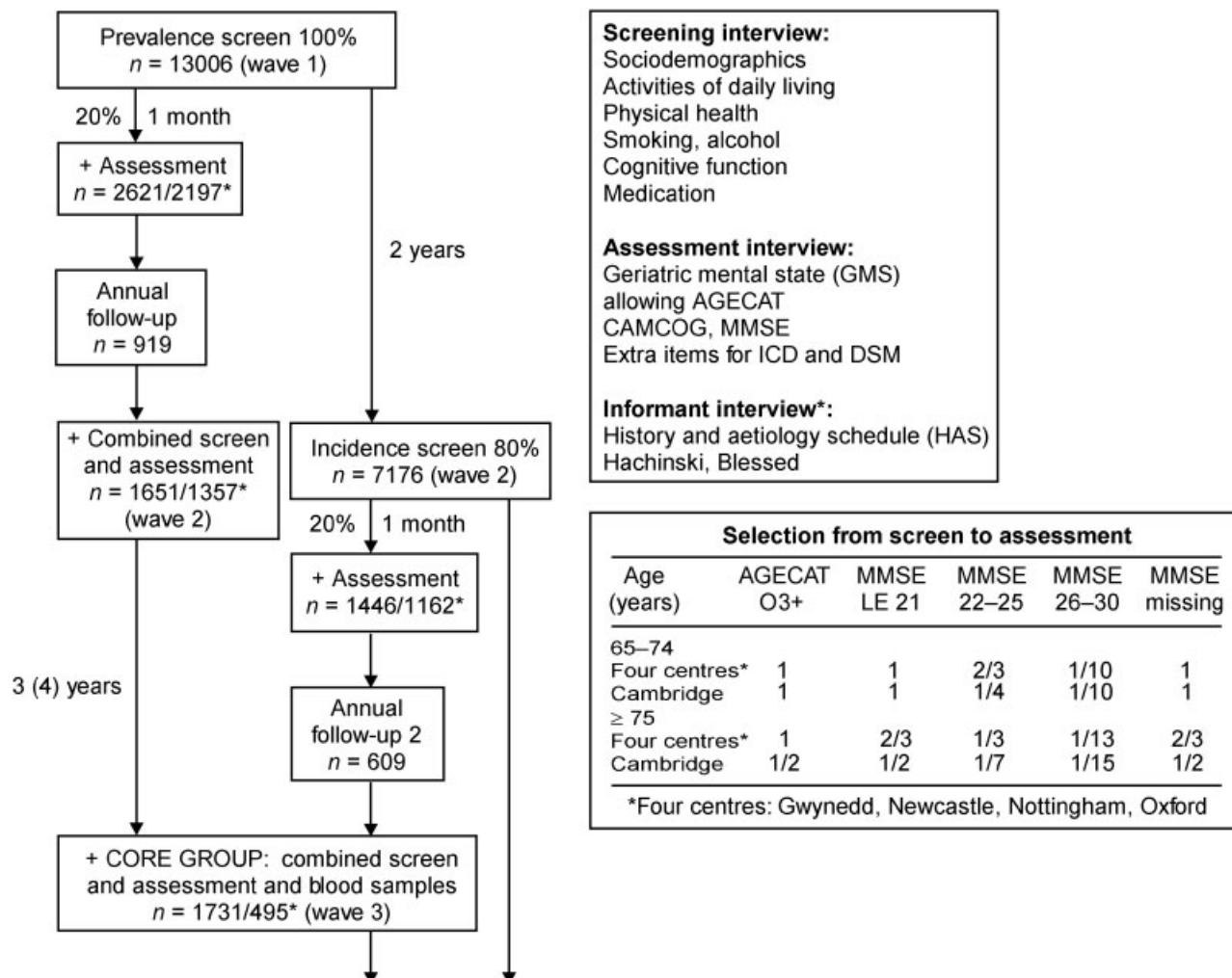


Figure 1. MRC CFAS: methodology (five identical centres). + Declaration of intent (DoI) to donate brain at post mortem. *Informant interview

centre (Liverpool) with a different design and funded earlier. Each of the five centres obtained a stratified random sample from Family Health Service Authority lists of sufficient individuals aged 65 years and over to achieve at least 2500 interviews (see Figure 1 for the study design). The Liverpool study consisted of a sample of 6035 individuals aged 65 years and over, stratified by sex and 5 year age bands. 5222 received a detailed assessment interview, with a selection also receiving the same interview by a clinician 3 months later. This process was repeated at 2 and 4 years. At 7–8 years the cohort was reinterviewed using the five-centre combined screen and assessment interview.

BRAIN DONATION

At all waves of the study, selected participants and their families have been approached with a request to consider donating brain tissue after death. Procedures are in place to collect, process and examine brains from these individuals, which are stored locally. There are currently 347 brains within

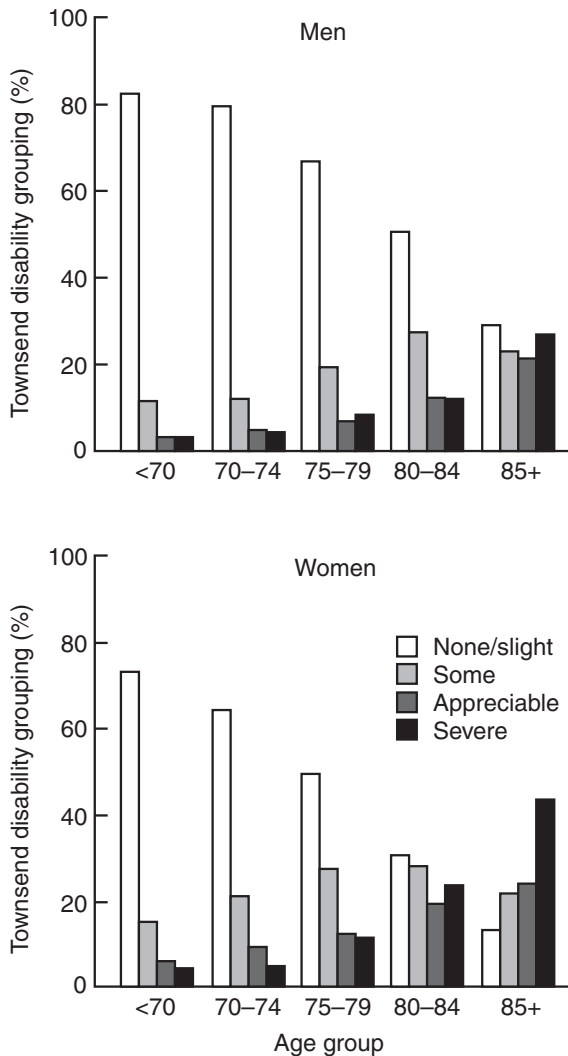


Figure 2. Percentage within Townsend disability grouping by age group and sex. From reference 2, with permission

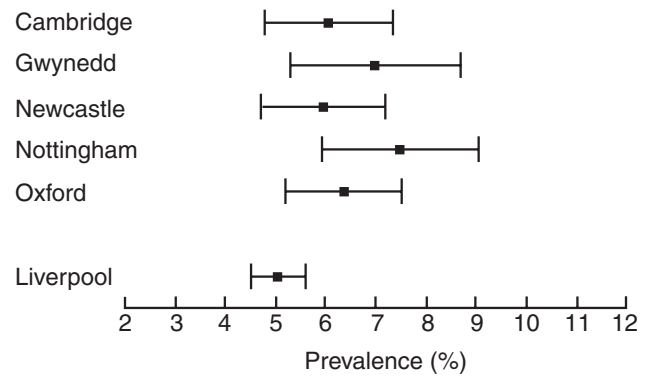


Figure 3. Prevalence estimates with 95% confidence intervals for AGE-CAT organicity O3+. From reference 3, with permission

Table 1. Estimated numbers of cognitively impaired or disabled elderly people in England and Wales by age group, sex and type of disability

Disability	64–74 years		75–84 years		≥85 years	
	Men	Women	Men	Women	Men	Women
Physical only	68 000	122 000	82 000	257 000	43 000	232 000
Cognitive only	32 000	29 000	45 000	57 000	20 000	61 000
Combined	14 000	11 000	26 000	57 000	27 000	99 000

From reference 1, with permission.

the Neuropathology study and 209 have been used for a first analysis. There are a further 500 people who have made a declaration of intent to donate brain tissue (DoI) and the continuing mechanism for collection will enable us to reach our target of 450.

BLOOD RESOURCE

During wave 2 in Oxford, wave 3 in Cambridge, Gwynedd, Newcastle and Nottingham and wave 4 in Liverpool, a blood sample (or saliva when blood was refused) was requested. There are 1126 blood and 193 saliva samples from the assessed groups in five sites (Cambridge, Liverpool, Gwynedd, Newcastle and Nottingham) and a further 1600 from the full wave 2 population in Oxford.

MORTALITY DATA

The full sample of 24066 is flagged on the NHS Central Register at the Office for National Statistics (ONS); 11 104 death notifications, with causes coded to ICD-9, have been received. The death information complements and enhances data from interviews, as all eligible for entry into the study, together with those actually interviewed, were flagged. This enables tracking of all individuals from initial sampling to death.

OUTPUT FROM THE STUDY

Figures 2 and 3 and Table 1 provide some results from published papers. A full bibliography may be found on the CFAS website, <http://mrc-bsu.cam.ac.uk/cfas>. Work in progress includes measures of healthy life expectancy, estimation of incidence rates, medication usage, normative values for CAMCOG and its subscales, cognitive function as a marker for survival, neuropathology analysis and genetic risk for dementia.

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The Epidemiology of Alzheimer's Disease: An Update

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This chapter provides a short overview on methodologic issues related to the design of epidemiologic studies of dementia, and provides an update on the epidemiology of Alzheimer's disease (AD). AD is characterized by a gradual but steady decline in cognitive and occupational function. A clinical diagnosis of AD is based on the course of the dementia and exclusion of other known causes of dementia. A definite diagnosis of AD can only be made post mortem in the presence of neuritic plaques and neurofibrillary tangles¹. Although these lesions are considered to be pathognomonic for the disease, it is unclear as to whether they are the primary cause of the disease or are the consequence of other more fundamental processes.

METHODOLOGY

Study Design

Research into the frequency and etiology of dementia is based on studies of prevalent cases and incident cases. Prevalent cases are identified in a cross-sectional study. The probability that they are found depends on the likelihood that they developed into a case and survived until the time of the survey. Thus, there is usually over-representation of cases with long duration. Incident cases are newly developed cases identified in a prospective follow-up study of a cohort that is initially dementia-free at baseline. It is preferable to study etiologic factors related to dementia in incident cases, as there is less likelihood that factors related to survival (or severity of the disease) influence the association of the exposure to the disease.

Measurement of Risk Factors

Ascertainment and interpretation of risk factor data is also less subject to bias when collected from incident, compared to prevalent, cases. In incident cases, a risk factor can be measured at baseline before the onset of dementia. In a prevalent case, a proxy needs to be questioned because a demented person can no longer be expected to provide valid or reliable answers. In addition, some biologic markers in prevalent cases may be influenced by the dementia itself².

Case Identification

Most cases of dementia do not come to the attention of the health services. This is the result of many factors, including the cases themselves or their caretakers not recognizing the disease³ and there presently being few treatment alternatives. Likewise, dementia is under-reported as cause of death. Therefore, studies relying on health care systems or records to identify cases probably only capture moderate to severe cases or cases with exceptional presentation. To fully identify the range of case presentation in the population, including mild cases, population-based studies that interview individuals in person are needed.

Diagnostic Guidelines

Strides in standardization of diagnoses used in research studies have been made in the past decade. Currently, the diagnosis of AD is made on the basis of internationally agreed guidelines for dementia (i.e. DSM-IV⁴ and ICD-10⁵) and specific criteria for AD. The most widely used criteria for AD are from the NINCDS-ADRDA⁶. Although application of different guidelines for dementia can identify different individual cases⁷, reasonable inter- and intra-rater reliability can be achieved⁸ if the same guidelines are used.

RISK FACTORS

Much progress has been made in the past decade in testing hypotheses based on case-control studies⁹, and also in identifying new hypotheses.

Age and Sex

All epidemiologic studies show an increase in the frequency of dementia with age, with an almost doubling of prevalence and incidence from age 65–85 years^{10,11}. Approximately 65% of cases are attributed to AD, although the proportion of AD cases out of the total might depend on ethnic group¹². Furthermore, as more neuroimaging is used in diagnosis, more cases of dementia associated with cerebrovascular

disease may be identified¹³. The slope of the increase with age depends on the sub-type of dementia. The incidence of dementia and AD increases steeply with age. The absolute incidence and the age-related increase in vascular dementia is lower. Studies differ concerning the sex-specific risk for AD. After 85 years of age a large difference in incidence was found in European studies¹⁴; the incidence in women increased steadily with age (up to 82/1000 person-years at 90 years and older), but flattened out in men (at 25/1000 person-years at 90 years and older). In contrast, in USA-based cohorts^{15,16}, the incidence of dementia, or AD, increased in old age with no apparent differences by sex. Sex differences in the risk for AD may be due to differences in biology, cumulative survival, or behaviour and exposures. Further investigations are needed to clarify the contribution of sex to the risk for AD.

Education

The contribution of education to the risk for AD is still controversial. Some argue that an association reflects confounding by socioeconomic factors, or diagnostic bias due to poorer performance on neuropsychological tests by individuals with low education¹⁷. Others argue that education is a marker for biological capacity that modulates when a person reaches the threshold of clinical dementia¹⁸. Several studies based on prevalent cases show that low education is associated with an increased risk for AD¹⁹. Studies based on incident cases are inconsistent, with some showing no relation²⁰ and others showing a relationship²¹ of low education to increased risk for AD. In one study the increased risk associated with AD was confined to women²².

Head Trauma

Reports on the relation of AD to head trauma with unconsciousness are inconsistent. Most studies are based on prevalent cases, where a proxy has had to be asked about the case's history of head trauma. These studies have either shown no effect²³, an increased risk for AD only in men²⁴, or only in women with head trauma²⁵. In a study based on US war veterans with independently documented history of head trauma during the war, there was a two-fold increased risk for AD²⁶. One report suggested that head trauma is a risk factor in the presence of the apolipoprotein E*4 allele²⁷ but two other studies have failed to confirm this^{24,28}.

Cardiovascular Risk Factors

One new area that is being investigated is the relation to AD of cardiovascular disease and risk factors. Several direct and indirect mechanisms may explain such associations, including ischemia, hypoxia, hemodynamic factors and neurotransmitter metabolism²⁹. Studies have reported an increased risk for AD associated with subclinical measures of atherosclerosis³⁰ and elevated levels of blood pressure². Indicators of glucose metabolism, including glucose and insulin levels³ and diabetes³² have also been associated with an increased risk for AD, although not consistently³³. In addition, cardiovascular risk factors have also been shown to increase the risk for AD, including smoking^{34,35} and diet³⁶. Since this is a relatively new area of investigation, confirmation of these findings in other studies is needed.

Steroid Hormones

Estrogen may be linked to AD through several direct mechanisms related to amyloid processing, neurotransmitter metabolism, cerebral blood flow or through cardioprotective pathways³⁷. Epidemiologic studies based on prevalent cases are inconsistent, some show a positive effect³⁸, others not^{39,40}. Prospective studies suggest that estrogen replacement therapy is protective⁴¹⁻⁴⁴. However, these observational studies may be detecting an association of AD to healthy behaviour, as hormone replacement therapy users tend to be healthier⁴⁵. Studies examining estrogen effects in men, as well as clinical trials, are needed to confirm the association of increased estrogen levels to a reduced risk for AD.

NSAIDs

Based on finding remnants of inflammatory processes in neuropathologic material of AD brains, it has been hypothesized that anti-inflammatory medications may reduce the risk for AD⁴⁶. There have been many case-control studies based on prevalent cases⁴⁷. Although some of these studies show no effect, those that do provide estimates of as much as an 80% reduced risk⁴⁸. More recently, several prospective studies have been published, with inconsistent results. Studies with one measure at baseline and subsequent follow-up found no association^{49,50}, and another showed a non-significant reduced risk among those using NSAIDs for 6 months or more within 10 years prior to the diagnosis of dementia⁵¹. In the Baltimore Longitudinal Study of Aging, the risk of AD decreased with increasing duration of NSAID use over a 16 year follow-up⁵². As with the estrogen hypothesis, that pertaining to NSAIDs needs to be tested in controlled clinical trials.

Genetics

With time, increasingly more will be known about the genetics of AD. Currently, several specific mutations have been found in familial (early onset) cases of AD, including missense mutations in the β -amyloid precursor protein on chromosome 21⁵³ and mutations in the presenilin-1 (chromosome 14q)⁵⁴ and presenilin-2 (chromosome 1) genes⁵⁵. However, these mutations do not account for the vast majority of cases, which are sporadic and of later onset. To date, apolipoprotein is the only identified polymorphism that has consistently been shown to be associated with a genetic susceptibility to sporadic and late-age onset AD. Specifically, the e*4 allele increases the risk for AD⁵⁶. Genes involved in inflammatory, apoptotic, metabolic, cytoskeletal, and neurotransmission processes are also under investigation⁵⁷. However, their role in modulating the risk for AD needs further elucidation. There are examples where a polymorphism is found to increase the risk for AD in one sample, but the findings cannot be replicated in others (i.e. such as the polymorphism in the gene encoding α 2-macroglobulin^{58,59}). Identification of genetic factors contributing to AD will not be simple. The majority of cases are probably a pool of heterogeneous conditions. It is not likely that there will be one or two major genes that are identified; rather, the genetic risk for AD will likely be the product of a number of genes that make a small contribution to risk.

SUMMARY

In the past decade many more prospective population-based studies were started and have yielded valuable data. However,

besides age and the apolipoprotein gene, few risk factors have been shown consistently to reduce, or to increase, the risk for AD. With advances in our understanding of the brain and our genetic code, rapid progress should be made in identifying environmental and genetic risk factors and in studying the interaction between the two.

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The Lundby Study, 1947–1997

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The Lundby Study is a prospective, longitudinal investigation of all kinds of mental disorders in a total population. It started in 1947, when four psychiatrists, Essen-Möller *et al.*, interviewed and described all the 2550 inhabitants of a geographically delimited area in southern Sweden. They were interested in finding out how many people were suffering from a mental illness or deviation, and also the way normal and abnormal personality traits were distributed in a normal population. They traced and examined all but 1% of the population. Only four people refused to participate. After 10 years, in 1957, the field investigation was repeated. One psychiatrist alone (Hagnell) re-examined the original population from 1947, irrespective of domicile. For deceased persons, information was collected from hospital case notes and from relatives. Hagnell also examined the 1013 newcomers in Lundby and thus performed a new prevalence study of 2612 people, as well as a 10 year incidence study of the original 2550 subjects. In 1972 Hagnell and Öjesjö made a second follow-up study¹ including the total population of 3563, also this time irrespective of domicile, thus performing a 25 year and a 15 year incidence study. The participation rate was as high as 98% at the two follow-ups.

All the three field examinations were performed in a similar way. One part of the examination was a semi-structured interview, the other part a free conversation. The task of the psychiatrist was to collect the information to the best of his ability, observantly looking for symptoms and signs. In an investigation such as this, the data should ideally be collected by a trained psychiatrist. To facilitate his task, the psychiatrist had a structured form to fill in. Each interview turned out to be different, but the information gathered was noted in a systematic way. Supplementary information was collected

from, amongst other sources, official registers, hospital case notes, autopsy reports, social insurance offices, local officials and relatives. This additional information from other sources forms a rich resource and has proved to be particularly valuable when evaluating the dementias of the elderly. All the types of information collected formed the basis for the global evaluation and classification.

The material from the three field studies has enabled us not only to estimate the rates of prevalence, incidence and probability of developing a mental illness, but also to identify possible background factors of, for example, dementia, starting with people who were healthy at one investigation but who had contracted the illness before the next. Trying to identify risk factors is a weighty enterprise in which we are still involved.

In an epidemiological, psychiatric investigation, precise clinical classification tools are hardly applicable. For dementias among the elderly we use two diagnostic groups: senile dementia of the Alzheimer type (SDAT) and vascular dementia (MID), the latter characterized by focal brain symptoms. Of course, other dementias of the elderly exist but they are fairly unusual compared with SDAT and vascular dementia and not very easily recognized in epidemiological investigations.

In order also to be able to include the hereditary aspect, we have mapped out the kinship of every proband down to the sixth generation, and for the original 2550 even down to the seventh generation.

From the Lundby Study, several reports on dementias in the elderly have been published. Figures from the 1972 prevalence study are included in the EURODEM comparative prevalence studies^{2,3}. For the 10 year age groups 60, 70 and 80, the prevalence

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rates of SDAT in the Lundby Study were, for both sexes taken together, 0.3%, 2.5%, and 10.9%, respectively, and for vascular dementia 0.8%, 3.5%, and 6.3%.

The rates of incidence and cumulative probability of contracting SDAT or vascular dementia were also published^{4,5}. The cumulative lifetime probability of contracting SDAT was reported to be 25.5% in men and 31.9% in women. If only the most severely impaired cases are counted, the figures are 15.0% and 22.2%, respectively. For vascular dementia the figures are 29.8% in men and 25.1% in women, for severe cases 16.6% and 15.2%, respectively.

Attempts have been made at finding background factors of a precipitating or protective nature in SDAT and vascular dementia in the Lundby Study. Two articles on this topic have been written^{6,7}.

On 1 July 1997 the Lundby Study^{1,8,9} celebrated its 50th anniversary by launching a new re-examination of its population, now aged 40+. This fourth wave of the Lundby Study gives an opportunity to study the occurrence of mental disorders and, as the population has grown 25 years older since 1972, especially mental disorders in an ageing population.

Between 1 July 1947 and 1 July 1972, 736 probands had died, and between 1 July 1972 and 1 July 1997, 1028 probands had died. On 1 July 1997, in all, 1764 probands were deceased. Thus, the number of living probands on 1 July 1997 was 1799 (3563–1764). This field investigation is now completed and the participation rate is still high (approximately 90%).

In the 1997 Lundby Study, modern diagnostic systems, such as DSM-IV¹⁰ and ICD-10¹¹, are used, together with the old diagnostic Lundby classifications. When the interview is finished, the proband is given a set of self-rating questionnaires. The proband is asked to complete them and to return them to the Lundby Study. The self-rating questionnaires are: the Interview Schedule for Social Interaction about social network¹², the Nottingham Health Profile about quality of life¹³, the Sense of Coherence Scale (about salutogenesis)¹⁴, and the Hopkins Symptom Check List, a mental symptom check list¹⁵. Also, this time supplementary information will be collected.

Focusing on the mental health/illness and its consequences as the present prospective, longitudinal study can shed light on the incidence and course of mental disorders in the middle-aged and elderly and estimate the importance of health care and treatment. By revisiting and examining the Lundby population a fourth time, the Lundby database will be updated. No comparable study has been progressing for such a long period of time.

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Nutritional Factors in Dementia

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The prevalence of nutritional disorders among the elderly is substantial, particularly among the elderly mentally ill (see Chapter 138a). Those with dementing disorders are at special risk. Dementia caused by a single nutrient deficiency is rare in the UK and USA and malnourishment will normally be the result of a dementia and deranged eating patterns, rather than its cause. The elderly in hospital or residential care are more likely to be malnourished than those living in the community^{1,2}.

Reduced plasma and blood levels of folate³⁻⁶, vitamin B₁₂³, vitamin C^{2,5,7-10}, and vitamin E⁸⁻¹⁰ in particular, have been reported in association with dementia and cognitive impairment.

Goodwin *et al.*¹¹ reported a direct relationship between water-soluble vitamins and cognitive function in healthy subjects. Those with the lowest levels of plasma folate and vitamin B₁₂ scored significantly worse on the Halstead-Reitan categories test and the Wechsler memory test. As part of the Basle Longitudinal Project on Ageing, involving urban-dwelling healthy elderly people, Perrig *et al.*¹² found that β -carotene and vitamin C levels were independently correlated with memory function after controlling for compounding variables such as age, education and gender. High intake of monounsaturated fatty acids was found to be associated with better memory function in 300 people aged 65–84 years¹³. Unsaturated fatty acids appear important for maintaining the development and integrity of neuronal function.

Botez *et al.*¹⁴ having found evidence of organic brain damage on psychometric testing in folate deficient subjects, reported improvement following 12 months' supplementation. Of five patients who underwent radionuclide cysternography, three showed improvement of cerebral atrophy. Supplementation with vitamin E has been associated with slower development of the pathology of Alzheimer's disease¹⁵.

It is not easy to establish, from studies of this sort, whether nutritional deficiency is a cause or effect of cognitive impairment. Supplementation studies suggest that certain additional nutrients may influence cognitive performance and, whether causative or not, nutrient deficiency may increase cognitive decline. If aetiologically significant, then the relevance of many of these findings may lie in the relationship with the oxidative stress hypothesis of Alzheimer's disease and the significance of oxidative stress for the development of ischaemic vascular disease. There is increasing evidence that reactive oxidizing species contribute to the neuronal damage and formation of the amyloid plaques seen in Alzheimer's disease^{16,17}. Antioxidants could be important for preventing this damage and antioxidant constituents of diet, such as vitamins E and C and β -carotene may be of particular importance. Vitamin C is actively concentrated in the brain¹⁸ and is considered to be the most effective antioxidant in human plasma¹⁹.

Impaired antioxidant status²⁰ and low plasma vitamin C levels²¹ have been suggested as risk factors for coronary artery disease and, therefore, may be relevant to dementia of vascular origin. Similar associations have been reported with vitamin B₆ and folate consumption²². Preliminary suggestions that vitamin E may reduce the risk of stroke has obvious implications for the development of vascular dementia.

The causative role of thiamine deficiency for the Wernicke–Korsakoff syndrome is established and it seems that some alcoholic dementias may result from Wernicke–Korsakoff lesions²³. Other B vitamins are capable of producing cognitive impairment and dementia²⁴ but are only likely to be significant causes in industrially underdeveloped countries.

Assessment of nutritional status should be part of the general management of dementing patients and nutritional deficiency corrected whenever possible. The cause of undernutrition is multifactorial and various interventions may help to improve a patient's nutritional state²⁵ (see Chapter 138a). For a few patients this may lead to gratifying improvements of cognitive performance, but all are likely to benefit by a reduction of morbid risk and improvements in general health. It is likely that a clearer understanding of the relationship between dietary constituents and dementia will contribute to methods of prevention and treatment of this condition.

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The Genetics of Alzheimer's Disease

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Knowledge of the genetics of Alzheimer's disease (AD) has advanced in recent years. Like many other diseases of late life, AD exhibits a complex pattern of inheritance. It is difficult to localize genes that cause such diseases, however, partly because only recent generations are available to be genotyped and phenotyped. This chapter will provide a general overview of the current state of knowledge in the genetics of AD and related dementias, and suggest approaches to interpretation of future findings in this area.

FAMILY AND TWIN STUDIES

The striking early evidence for genetic causes of AD was based on a few multigenerational kindreds with many individuals with the disease (for review, *see ref. 1*). Most of these families had onset of disease before the age of 65, and were thus arbitrarily classified as having "early onset". These multigenerational kindreds with so-called familial AD probably represent no more than 1–2% of all AD cases, but their existence strongly suggested that genes were important in at least some cases of AD.

Other early work supporting a genetic etiology of AD was based on the presence of a positive family history of the disease in clinical and population-based AD samples. Most of these studies showed an increased risk of AD in first-degree relatives, ranging from 25% to 50%, as compared with 10% or less for non-demented controls². This finding suggested an up to five-fold increase in risk of AD among relatives of individuals with the disease.

The early family studies focused on families at greater risk of developing AD. Recently, however, some researchers have taken another approach to the problem by investigating families who appear to have a *lower* risk of AD³. They have shown that the first-degree relatives of individuals who reached the age of 90 and were not demented had a reduced liability to AD from age 60 on, as compared with the families of younger non-demented individuals. The message here is that genes may either promote or reduce the risk of AD.

Twin studies have provided other evidence for genetic causes of AD. Members of identical (monozygotic) twin pairs share 100% of their genome, while fraternal (dizygotic) twin pairs share 50% of their genes on average. The twin design assumes that the degree of similarity in the early life environments of identical and fraternal twin pairs are the same. Based on this so-called "equal environments assumption", the twin design compares concordance rates for disease in identical (monozygotic) twin pairs with that in fraternal (dizygotic) pairs. Higher concordance in identical pairs, then, implies genetic influences. The magnitude of such influences can be estimated in complex models that simultaneously assess the proportion of disease liability attributable to

shared or unique environmental causes, as well as stochastic variation⁴.

Like the early family studies, twin studies of AD have generally implied genetic causation without identifying particular genes or loci. The heritability of AD (proportion of disease liability attributable to genes) has been estimated from ongoing studies of AD in three population-based twin registries: (a) the NAS–NRC Twin Registry in the USA⁵; (b) the Swedish Twin Registry⁶; and (c) the Norwegian Twin Registry⁷. Based on tetrachoric correlation analyses, current estimates of heritability for AD in these studies range from 0.33 to 0.74^{5–7}. We suggest that the wide range of these estimates of heritability may reflect the differences in the age distribution of twins in the three registries. We speculate that the lowest heritability estimate (0.33) from our work may underestimate the true heritability of AD because of the atypically young age (mean onset age = 67 years) of AD cases in the NAS–NRC sample. Others have shown that the risk of developing AD, for a relative of a proband with AD, increases with age at least until the mid-80s⁸. Consistent with this, we have found that the estimates of heritability in the NAS–NRC Registry have gradually increased over the past few years. As the Registry approaches the typical age of risk for AD of the late 70s, 80s and even beyond, we anticipate that heritability in the NAS–NRC sample will approximate that of the other two studies and the consensus estimate of heritability will likely be between 0.5 and 0.65.

MOLECULAR GENETICS

Autosomal genes with dominant expression

Motivated by the findings from family and twin studies, molecular genetic techniques have now identified mutations at three genetic loci that are associated primarily with early onset AD. Initial interest in the molecular genetics of AD focused on chromosome 21, because aging Down's syndrome patients (trisomy 21) often exhibit the brain pathology seen in AD patients^{9,10}. The first identified AD mutation was found in the β -amyloid precursor protein (*APP*) located on chromosome 21¹¹. This mutation substitutes isoleucine for valine at codon 717 in exon 17¹¹. Since the first report of this missense mutation, a number of other *APP* mutations have been identified (for review, *see ref. 12*). But, in total, the *APP* mutations appear to account for the disease in less than 20 families worldwide.

The second genetic locus to show linkage to AD was found on chromosome 14^{13,14} and was termed presenilin-1 (*PS-1*)¹⁵. Over 40 mutations in the *PS-1* locus have now been reported in at least 82 families world-wide¹⁶. Like the *APP* mutations, *PS-1* mutations

appear to act as autosomal (i.e. not sex-linked) dominant traits with nearly complete penetrance¹⁶. Early reports suggested that chromosome 14 mutations may account for the majority of early-onset familial AD cases^{17,18}. More recent estimates from population-based samples^{16,19} have produced conflicting estimates of the proportion of early-onset cases attributable to *PS-1* mutations, with some¹⁶ suggesting that *PS-1* mutations account for less than 20% of such cases (see p. 217 for more discussion of presenilins).

A third autosomal dominant locus for AD has been localized to chromosome 1 and termed presenilin 2 (*PS-2*)^{20,21}. To date, *PS-2* mutations have been identified in only a few families of different ethnicity. In fact, in a population-based sample of early-onset AD cases, it was estimated that *PS-2* mutations account for less than 1% of the cases¹⁶. The *PS-2* mutations usually provoke onset of AD symptoms before age 65, but later onsets occasionally occur. The variability in onset age for the *PS-1* and *PS-2* mutations suggests the complexity of these otherwise Mendelian traits.

Identification of AD mutations on chromosomes 21, 14 and 1 has led to major advances in understanding the etiology of this disease. It is important to note, however, that these mutations probably account in total for less than 2% of all AD cases²². Importantly, it appears that they do not account even for the majority of so-called autosomal dominant familial AD¹⁶. Thus, there are almost certainly other major genes remaining to be identified in early-onset AD. A separate and, from a public health perspective, far more compelling issue is the search for other genetic influences in common, late-onset AD.

Apolipoprotein E

In the search for other so-called AD genes, Pericak-Vance and colleagues²³ reported linkage of disease to a locus on chromosome 19 in pedigrees with late-onset AD. Searching through the candidate genes of the implicated region on chromosome 19, Strittmatter *et al.*²⁴ found that the apolipoprotein E gene (*APOE*), which encodes the lipid transporter apolipoprotein E (apoE), is located in the same region that showed linkage to familial late onset AD. Their initial studies showed an increase in amyloid deposition in elderly individuals with the $\epsilon 4$ allele at *APOE*^{24,25}, and an increased frequency of this allele in both familial and sporadic AD cases²⁵⁻²⁹. Subsequent studies showed that, as the number of $\epsilon 4$ alleles increases from 0 to 2, the age of onset of AD decreases³⁰. These findings have now been confirmed in over 100 studies, including some that extend the findings to early onset AD (for review, see ref. 31). A recent meta-analysis of data on over 14 000 subjects from 40 research teams synthesized the findings on *APOE* through 1997³². That study confirmed the basic findings noted above and also replicated some findings that previously had been suggested by select studies. These findings were that the *APOE* $\epsilon 2$ allele may be associated with lower risk of AD, and that risk associated with the $\epsilon 4$ allele appears to vary by age, sex, and ethnicity. The association between AD and *APOE* $\epsilon 4$ is complex, and it is not clear why some individuals with the $\epsilon 4$ allele develop the disease while others do not. But, whatever its precise role, $\epsilon 4$ is an important player in the genetics of AD, as various groups have now estimated that *APOE* accounts for 57–70% of the genetic contribution of AD³³⁻³⁵. Further discussion of the role of *APOE* on the risk of AD is provided on pp. 218–19.

Other Putative Genes

There are many pedigrees that show intense familial aggregation of either early- or late-onset AD but have no identified mutation and no *APOE* $\epsilon 4$ allele. It seems likely, therefore, that one or more genes predisposing to AD remain to be discovered. The pursuit of

these other genes has produced numerous reports of associations between specific genes and AD, but none of these has been consistently replicated. Chromosome 12 has been the focal point of much of this work, since a genomic screen showed a region on this chromosome may be associated with increased susceptibility to AD in individuals without an *APOE* $\epsilon 4$ allele³⁶. Further analyses by this group has suggested that a region on chromosome 12 near the one of original interest may be associated with increased risk of the Lewy body variant of AD³⁷. Other groups have now demonstrated linkage to regions near the initially implicated chromosome 12 locus, although one such study suggested that the strongest such association was in individuals with an $\epsilon 4$ allele³⁸, while the other showed the strongest association in those without an $\epsilon 4$ allele³⁹.

Two of the several candidate genes in the region under scrutiny on chromosome 12 have now been shown to be positively associated with AD in at least some studies. The first is the gene for the major apoE receptor in the brain, the low-density lipoprotein receptor-related protein (LRP) that is selectively found in neurons and reactive astrocytes⁴⁰. This gene, *LRP*, has also been shown to mediate the endocytosis and degradation of *A β* ⁴¹. It has two alleles, called C and T. The homozygous C genotype appears to be associated with increased risk of AD, earlier onset of AD, and significantly more neuritic plaques at post mortem, as compared to individuals with at least one T allele⁴². Several groups have confirmed these results⁴³⁻⁴⁶, but at least two have not^{47,48}.

The other candidate locus encodes $\alpha 2$ macroglobulin (*A2M*) and is located just outside the area on chromosome 12 for which others have reported linkage³⁶. Previous work⁴¹ showed that *A2M* is a major LRP ligand and is a serum pan-protease inhibitor that mediates the clearance and degradation of *A β* , the major component of brain amyloid. At least three different *A2M* polymorphisms have been implicated so far⁴⁹⁻⁵¹. One of these studies reported that, in a very elderly Finnish cohort, the association with the *A2M* polymorphism was only evident in neuropathologically diagnosed AD cases, but not in the clinically diagnosed AD cases. This particular polymorphism was also associated with an increase in the neocortical β -amyloid protein load⁵¹. Unfortunately, a number of other studies have not confirmed these associations between *A2M* and AD in several large samples⁵²⁻⁵⁶. Together, these findings suggest that if *A2M* is associated with AD, its effect may be limited to subgroups of susceptible individuals and that different polymorphisms may be influential in each of these subgroups.

With so many purported loci and mutations associated with AD, one may wonder about a common thread in the pathogenesis of AD. The pathogenic mechanisms of the *APP*, *PS-1* and *PS-2* mutations are not completely understood, but each appears to be associated with increased production of the long form of *A β* (*A β* -42), relative to the production of the shorter forms (mostly *A β* -40) of *A β* ⁵⁷. *A β* -42 seems to be a particularly pathogenic form of *A β* in AD. Both *LRP* and *A2M* are involved in the degradation of *A β* . It has been suggested that these links to *A β* may be the underlying common pathogenic event leading to AD for these genes⁵⁸.

There are reports of numerous other genes throughout the genome contributing to AD, but none of these findings has been consistently confirmed. A partial list of the contending genes includes: $\alpha 1$ antichymotrypsin (*ACT*), a gene on chromosome 14⁵⁹⁻⁶⁷; the K variant of butyrylcholinesterase (*BCHE-K*) on chromosome 3⁶⁸⁻⁷²; bleomycin hydrolase (*BH*) on chromosome 17⁷³ (cf. ref 74); the non-amyloid component precursor gene (*NACP*/ α -synuclein) on chromosome 4⁷⁵ (cf. ref 76); the human leukocyte antigen (*HLA*) genes on chromosome 6^{77,78}; the *FE65* gene on chromosome 11⁷⁹; the dihydrolipoyl succinyltransferase (*DLST*) gene on chromosome 14^{80,81}; the

interleukin-1 (IL-1) on chromosome 2^{82,83}; and the mitochondrial cytochrome *c*-oxidase (CO) genes *CO1* and *CO2*⁸⁴. To make matters more complicated, many of the above-named genes have been shown to interact in varying degrees with *APOE* or other genes to alter risk of AD⁸⁵.

There are several possible reasons for the contradictory results of the association studies noted above. Other authors have proposed that they may reflect: (a) different etiologies in different ethnic populations; (b) lack of adjustment for multiple comparisons (type I statistical error, resulting from simultaneous screening of many different candidate loci); (c) linkage disequilibrium in some (or all) populations between the tested polymorphism and the functional polymorphism; or (d) inability to detect an effect due to limited statistical power or weak genetic effects^{39,86}. Association studies are also more difficult to interpret when one does not know such characteristics as the mode of inheritance, age-dependent expression characteristics, interaction with other loci, misdiagnosis rate, and disease allele frequency. Finally, the inconsistent findings may reflect interactions with non-genetic factors. There is convincing evidence implicating *non-genetic* factors in the etiology of AD, either alone or in interaction with specific genes^{87,88}. The most compelling indications of this effect are the numerous reports of monozygotic twin pairs who, although genetically identical, remain discordant for AD for 10 or more years^{5,89-91}.

Collectively, the three twin studies previously described suggest that non-genetic influences account for at least 25% of the variance in AD susceptibility. Studies of these "environmental factors" have suggested that both potentiating and protective influences may operate by primarily altering the timing of onset⁹²⁻⁹⁴. Similar to the recent reports of genetic associations, however, few non-genetic risk factors have been consistently confirmed. For further discussion on the genetic epidemiology of AD, see pages 219-21.

CONCLUSION

Given the number of AD genes already identified and the proportion of cases attributed to each of these genes, we can reasonably predict that there will be more than one other new gene implicated in AD. Based on what we know about the complexity of the relationship of AD and the genes identified to date, it is likely that any newly identified genes will also interact with other genes and with non-genetic factors. Also, the effect of any one gene may be influenced by the age, sex, and ethnicity of the at-risk individual. Therefore, any newly identified genes will need extensive evaluation and confirmation in various subpopulations before they are added to the list of AD genes.

Until the search for genetic and non-genetic causes of AD is complete, there is no way of identifying those individuals who will definitely develop AD. However, the discovery of new AD genes may have benefits beyond just identifying those at increased genetic risk for the disease. At least two lines of evidence suggest that there may be a common underlying pathological mechanism regardless of the genes involved. First, multiple predisposing genes appear to produce the same AD phenotype. Second, all of the AD genes identified to date, and some of the other candidate genes under examination, have some link to the metabolism of $A\beta$, a neuropathological marker of the disease. Understanding this common feature may lead to evidence of a common destructive metabolic process (i.e. final common pathway) and may be the key to identifying susceptible individuals *before* they develop the full syndrome of AD, or even preventing the disease. Because clinically normal individuals with at least one *APOE* $\epsilon 4$ allele have "AD neuropathology" as early as the fourth decade of life⁹⁵, one might postulate that other "AD genes" may also begin

their destructive process early in life. If one could find the common pathological mechanism for the disease, then regardless of the genetic or non-genetic factors that initiate the process, treatment could be implemented to delay or prevent progression of the disease. The underlying premise is that it may be more feasible to identify a "marker for the disease process" and develop a test to measure a single marker present fairly early in life, rather than try to develop an algorithm to predict AD that includes all possible variations of genetic and non-genetic risk factors for disease.

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PS-1 protein occurs within the perikaryon and dendrites of nerve cells, mostly within the early Golgi apparatus and smooth endoplasmic reticulum. No difference in distribution or amount of

PS-1 protein or its message are seen in PS-1 AD. The holoprotein is processed into 17 kDa, C-terminal (CTF) and 27–28 kDa, N-terminal (NTF) fragments which accumulate in 1:1 stoichiometry, in a highly regulated and saturable manner. Cleavage occurs within the hydrophilic loop, around amino acid 298². PS-1 (2) mutations alter neither the site nor the manner of endoproteolysis, although the cleaved products may accumulate through increased stability.

Amyloid precursor protein (APP) trafficking from Golgi to endosome and plasma membrane—the principal sites of A β production—is not affected by PS-1 mutations. PS-1 knockout mice show normal α - and β -secretase activity, though A β production is decreased with accumulation of the C-terminal stub of APP. PS-1 therefore might regulate γ -secretase, as part of the γ -secretase complex, with mutant PS-1 fragments enhancing this activity^{3,4}.

CTF and NTF of PS-1 form a stable complex containing GSK-3 β and β -catenin⁵. The cellular trafficking, stability and turnover of β -catenin is altered by PS-1 mutations³ and these might interfere with the binding of GSK-3 β to tau, possibly promoting hyperphosphorylation and neurofibrillary tangle formation.

PS-1, therefore, probably regulates the sorting and processing of integral membrane proteins, including APP. PS-1 mutations may alter protein topology so as to favour APP catabolism along routes which, by increasing production and tissue deposition of A β , facilitate the pathological cascade. Other putative roles for presenilins could include signal transduction, involving *Notch* and *Wnt* pathways, during development or apoptosis⁴. Whether these functions are disturbed by PS-1 mutations, with repercussions for the pathogenesis of AD, is unknown.

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Apolipoprotein-E (APO-E)

Dan G. Blazer

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Strittmatter *et al.*¹ found that the gene apolipoprotein-E (APO-E) for the lipid transporter apolipoprotein-E is located on chromosome 19 in a region that was demonstrated to have strong linkage to familial late-onset Alzheimer's disease (AD). There are three common alleles formed at the polymorphic APO-E locus, E2, E3 and E4. Each individual inherits one allele from each parent; thus a person may have an APO-E genotype of E4/E4, E4/E3, E3/E3, and so forth². The E3 allele is the most common and represents approximately 78% of all alleles in European and American White populations. The E4 allele frequency is approximately 15–16% and the E2 frequency approximately 7%. The risk for AD is increased and the average age of onset is decreased with increasing numbers of APO-E E4 alleles. In addition to AD, the E4 allele has been shown to be associated with cardiovascular disease, renal disease, stroke, decreased ability to recover from physiologic challenges such as amnesia and an increase in all-cause mortality. Several case-control and incidence studies of late-onset AD have shown E4 allele frequency to be more frequent in cases; 30–50% in both sporadic and familial AD (FAD), compared to the 15% in the general population. The odds ratio for E4 allele heterozygotes developing AD are approximately 2–5, whereas the odds ratios for E4 homozygotes are 5–18. The E2 allele may actually retard the development of AD.

Other studies of AD and APOE in different ethnic groups report similar associations between the E4 allele and AD. Nevertheless, the frequency of the E4 allele among African-Americans appears to be higher than in Whites, yet the frequency of AD does not appear to be greater (thus suggesting less association, perhaps, in African-Americans compared to Whites).

Despite the association of AD and the APOE-4 allele, most persons who experience AD do not express the E4 allele. In addition, there have been reports of individuals homozygous for E4 in very late life who nevertheless remain cognitively intact. The E4 allele is therefore neither necessary nor sufficient to cause AD. For this reason, the E4 allele has been considered a susceptibility gene. This represents a change in perspective from the typical approach to genetic expression for an autosomal dominant or autosomal recessive.

Most genetic determinants of health do not derive from a one gene–one disease paradigm, but rather a paradigm in which the phenotypic expression of the genome is best conceived quantitatively. A single polymorphism at a specific locus can lead to multiple adverse outcomes, outcomes which can be investigated as changes over time as well as the onset of a specific disease at a specific point in time. In other words, these are susceptibility polymorphisms that are universally distributed in the population, rather than mutations that are uncommon and family-specific. A single allele may be sufficient to cause a specific disease, yet everyone with that allele has a measurable age-dependent risk for that disease. These susceptibility polymorphisms are therefore subject to investigation in the way epidemiologists conceive environmental stressors (such as stressful life events) or health-related behaviors (such as smoking), that is, as risk factors for disease onset and change. To date, however, few such susceptibility polymorphisms have been identified. Studies of these polymorphisms have focused almost exclusively upon the association of an allele with a specific disease or specific adverse outcome, such as mortality. Yet these polymorphisms may increase the risk

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Most genetic determinants of health do not derive from a one gene–one disease paradigm, but rather a paradigm in which the phenotypic expression of the genome is best conceived quantitatively. A single polymorphism at a specific locus can lead to multiple adverse outcomes, outcomes which can be investigated as changes over time as well as the onset of a specific disease at a specific point in time. In other words, these are susceptibility polymorphisms that are universally distributed in the population, rather than mutations that are uncommon and family-specific. A single allele may be sufficient to cause a specific disease, yet everyone with that allele has a measurable age-dependent risk for that disease. These susceptibility polymorphisms are therefore subject to investigation in the way epidemiologists conceive environmental stressors (such as stressful life events) or health-related behaviors (such as smoking), that is, as risk factors for disease onset and change. To date, however, few such susceptibility polymorphisms have been identified. Studies of these polymorphisms have focused almost exclusively upon the association of an allele with a specific disease or specific adverse outcome, such as mortality. Yet these polymorphisms may increase the risk

for multiple diseases, and therefore for generalized morbidity, and increased the risk of mortality from multiple causes. This translates into increased burden on the health care delivery system and higher costs.

Apolipoprotein E is a major serum lipoprotein involved in cholesterol metabolism. Lipoproteins derived from E4 are cleared more efficiently from the blood than those derived from E3 and E2. Apolipoprotein E does not cross the blood-brain barrier but is synthesized in the brain by astrocytes. In the brain, APOE is thought to be involved in the mobilization and redistribution of cholesterol and phospholipid during membrane remodeling associated with the plasticity of synapses³.

The biological basis for the association between the E4 allele and AD is unclear, however. APOE is found in senile plaques and fibrillary tangles and binds to A β in the cerebrospinal fluid. *In vitro* studies have indicated that APOE isoforms may differentially affect deposition of A β . Binding studies suggest that APOE isoforms have different affinities for A β . However, these results are controversial because different studies suggest different isoforms have the highest affinity for A β . The APOE isoforms have been shown to promote A β fibril formation *in vitro*. Other studies, however, show that E3 but not E4 binds to tau, preventing aggregation of tau and neurofibrillary tangle formation. Neurite extension and branching is more extensive in E3-treated cells compared with E4-treated cells. APOE E4 has also been shown to have increased antioxidant activity compared with E3³.

Although there is a consensus that APOE E4 is strongly associated with AD, this does not mean that tests for the genotype (which are commercially available) should be used for diagnostic testing. A consensus working group has recommended that, at the present time, APOE genotyping should not be used for predictive testings. APOE genotype alone does not provide sufficient sensitivity or specificity to allow genotyping to be used as a diagnostic test. Because AD develops in the absence of E4 and because many patients with APOE E4 seem to escape the disease, genotyping is also not recommended for use as a predictive genetic test⁴.

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Down's Syndrome and Alzheimer's Disease: Update

David W. K. Kay and Brian Moore

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One of the unresolved questions for the dementia of Down's syndrome (DS) as a model for Alzheimer's disease (AD) is the long interval, often up to 20 years, that separates the appearance of the AD pathology in the brains of individuals dying after the age of 35 from the clinical manifestations of dementia.

EPIDEMIOLOGY OF DEMENTIA

Study of decline and dementia is important for service as well as theoretical reasons. The life expectancy of a 1 year-old DS child with mild/moderate handicap is now 55 years². However, the wide variation in premorbid abilities and the presence of physical, particularly sensory impairments¹ often makes the diagnosis of dementia difficult. With ICD-10 criteria as standard, the Dementia Questionnaire for Persons with Mental Retardation³ and the Dementia Scale for Down Syndrome⁴ both perform well, but the Mini-Mental State Examination is too difficult⁵. Both caregiver information and longitudinal monitoring of cognitive performance are desirable⁶. Cortical atrophy may be demonstrated by brain imaging^{7,8}. Dementia is frequently associated with late-onset epilepsy, and late-onset epilepsy is associated with clinical evidence of dementia⁹⁻¹¹. Serial EEGs may reveal diffuse

abnormalities and slowing of the dominant rhythm associated with the decline of cortical functions¹².

In a meta-analysis of prevalence studies, the age distribution of dementia onset in DS was unimodal, with mean age of onset 51.7 years (SD 7.1, range 31-68), and earlier onset in women¹³. Age-specific prevalence rates of dementia in population-based samples were: age 30-39, 2.0-3.4%; age 40-49, 9.4-10.3%; age 50-59, 36.1-40%; and age 60-69, 54.5%¹⁴⁻¹⁶. DS subjects aged 50 years or over were significantly impaired on memory tests compared with younger subjects¹⁷. Dementia rates were increased in elderly people with learning disabilities (LDs) as a whole¹⁸ but the rates in DS were higher¹⁹. Persons with DS and dementia are also reported to show more non-cognitive symptoms than other persons with LDs and dementia²⁰. Dementia contributed significantly to decline in adaptive behaviour and skills after age 40, independently of age, while absence of medical illness had a favourable effect²¹.

Longitudinal Studies

Adaptive behaviour deteriorates more in DS than in matched controls with other LDs²². Treatable conditions, such as

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Longitudinal Studies

Adaptive behaviour deteriorates more in DS than in matched controls with other LDs²². Treatable conditions, such as

depression and medical illness, may be involved²³, but dementia is the main cause; in the absence of dementia and physical illness no decline was found²⁴. However, the incidence of cognitive decline and dementia seems to be very sensitive to the criteria used. Adults aged 22–56 who were dementia-free at baseline showed little change over a period of 3–4 years in most cases²⁵ and even over 6 years, persons aged 50 and over showed, on average, only very slight decreases in cognitive performance, attributed to precocious ageing; only 4% satisfied the criteria for dementia²⁶. However, on a sophisticated neuropsychological test battery, 28% of testable persons aged 30+ were deemed to show cognitive deterioration after 4 years, and of those aged over 50 only 30% showed no cognitive decline; as in AD, memory and learning were affected early²⁷. Whether persons with DS age prematurely or whether they age normally but are at increased risk of AD remains controversial. Further national and cross-national studies of the prevalence, incidence and natural history of dementia in both DS and in other LD are required²⁸.

NEUROPATHOLOGY AND GENETICS

The gene coding for familial Alzheimer's disease is now known to be distinct from the amyloid precursor protein (APP) gene, which is also situated on the long arm of chromosome 21. Overexpression of the triplicated APP gene results in higher concentration of APP in DS brains than in brains of elderly controls or of patients with AD²⁹. The increased amyloid deposition in AD could be due to some cases of AD being trisomy-21 mosaics, resulting from non-dysjunction during mitosis; such cases might possess other features of DS³⁰. A shared susceptibility to non-dysjunction and AD could account for an increased risk of dementia reported among the mothers of persons with DS³¹. Meta-analysis of seven studies of the ApoE polymorphism in DS found a similar distribution between DS adults and non-retarded controls, and no significant difference between DS persons with and without dementia³². There were, however, trends for the $\epsilon 2$ allele to be associated with later onset of dementia and later age at death in non-dementing persons, and for the $\epsilon 4$ allele to be associated with earlier onset. Cognitive decline in DS may be influenced by an unidentified gene situated at D21S11³³.

Clinico-pathological Correlations

The first change is the deposition of β -amyloid peptide ($\beta/A4$) in the cerebral cortex and elsewhere, in the form of diffuse, amorphous plaques, and may be seen as early as the second or third decade; soon afterwards microglial cells and ubiquitin protein can be detected, and by the age of 35 years "cored" amyloid deposits are seen at post mortem in most subjects³⁴. At this stage neurofibrillary tangles (NFT) are numerous in parts of the hippocampus and amygdala but are seen only in isolated cortical neurons. Eventually, and usually after the age of 50, the distribution and density of senile plaques and NFT and loss of neurons conform to the pattern seen in AD³⁴. This is the age when clinical dementia usually appears¹³. This gradual evolution of pathological change partly explains the puzzling temporal discrepancy between the onset of pathological and clinical manifestations of dementia in DS. Whether or not a similar evolution of neuropathological change occurs in AD is still unknown.

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International Criteria for Alzheimer's Disease and Their Problems— ICD-10, DSM-IV and NINCDS–ADRDA

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Dementia often comes to attention when an elderly person's cognition or behaviour no longer conforms to what is expected. Given inherent variability in people and their circumstances, in their past and present performance, in how this matters, in the expectation of changes with age, it is no surprise that how dementia is described, whether it is attributed to Alzheimer's disease (AD), and what that is taken to mean will vary within and across societies¹. Comparing three sets of commonly-used criteria for AD shows that while each conceptualizes dementia similarly, differences in their literal application can give highly variable results. Such spurious variability distracts from the important task of better understanding heterogeneity in disease expression—especially in the presence of medical and psychiatric co-morbidity.

THE THREE SETS OF CRITERIA

The criteria of the National Institute of Neurological and Communicative Disorders and Stroke, and the Alzheimer's Disease and Related Disorders Association (NINCDS–ADRDA)² require a clinical examination to produce evidence of dementia, defined as progressive deficits in memory and other areas of cognition (notably language, motor skills and perception) that occurs without a disturbance of consciousness, and in the absence of “systemic disorders other than brain diseases that in and of themselves could account for the deficits”. The criteria also require that the onset be between ages 40 and 90, the initial examination include a standardized cognitive test, and the deficits be “confirmed by neuropsychological tests”. “Exclusion of causes of dementia other than Alzheimer's” allows a diagnosis of *probable* AD, with neuropathological confirmation required to make the diagnosis *definitive*. *Possible* AD exists when “atypical” features or other co-morbid illnesses exist.

The 10th International Classification of Diseases (ICD-10) has both clinical guidelines³ and research criteria⁴ for Mental

and Behavioural Disorders. AD is classed as a dementia, defined as progressive impairment, in the absence of clouding of consciousness, of multiple higher cortical functions, specified as: memory, thinking, orientation, comprehension, calculation, learning capacity, language and judgement. Interestingly, the criteria also note that non-cognitive features (“deterioration in emotional control, social behaviour, or motivation”) can accompany or even precede dementia. The diagnosis is one of exclusion; only insidious onset and slow deterioration are cited as characteristic of AD. Subtypes include early and late onset, and “atypical or mixed”. The research criteria specify mild, moderate and severe stages, based chiefly on the extent of memory loss, although the impact on daily activities is noted.

The fourth edition of the *Diagnostic and Statistical Manual of Mental Disorders* (DSM-IV) of the American Psychiatric Association defines dementia as “multiple cognitive deficits that include memory impairment and at least one of . . . aphasia, apraxia, agnosia, or a disturbance in executive functioning”, provided that such deficits cause impairment, “occupational or social functioning” and represent a decline⁵. The criteria for “Dementia of the Alzheimer's Type” emphasize the need to rule out other conditions, and to exclude the syndrome if it exists only in the setting of delirium. Subtypes include early and late onset, and each of coincident delirium, delusions, and depressed mood. The accompanying text emphasizes insidious onset and gradual decline, notes that (but does not specify how) “cultural background” should be taken into account, and, while recognizing differences in levels of disability, provides no specific criteria for staging.

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Despite cross-national studies of their individual reliability and validity⁶ variability in diagnostic criteria can have profound

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PROBLEMS WITH CURRENT CRITERIA

Despite cross-national studies of their individual reliability and validity⁶ variability in diagnostic criteria can have profound

effects on resulting epidemiological⁷ and clinical⁸ estimates. This is especially true with strict exact adherence to each criterion, e.g. when patients whose dementia is too severe to sit for neuropsychological tests are said not to have fulfilled the NINCDS–ADRDA criterion that such tests be administered. Now while it might be argued that there is no point in criteria except in having strict adherence to them, we cannot ignore that most such criteria are revised periodically, and that the operationalization of some criteria (notably, functional impairment) can vary widely, again with marked impact on the resulting estimates⁹. A better approach would be to make clear how the interpretation of specific criteria may affect a study's results⁷.

Each set of criteria views AD as a diagnosis of exclusion. Given that AD is common and that many of the diagnoses to be excluded are rare, this approach seems perverse, despite the absence of a biological marker for the most common disorder. Perhaps it is the lack of a detailed consideration of staging, another problem with all the criteria, that has resulted in the “diagnosis of exclusion” approach. Without some understanding of systematic variation by stage, the heterogeneity of AD would seem chaotic, so that the clinician could only feel confident in the diagnosis when everything else had been rejected. By contrast, recognizing common features in how AD presents and progresses makes the task of diagnosing it with confidence much easier—indeed, where characteristic milestones are present, such confidence can extend even to a retrospective diagnosis¹⁰. This notion of a usual progression is, in fact, implicit in considerations of “atypical presentations”, which usually are understood as deficits (e.g. aphasia, apraxia) that occur at an *uncharacteristically early* stage. Also, in emphasizing insidious onset, the criteria seem to exclude patients whose dementia initially presents with delirium, despite that sequence being a common path to AD¹¹.

Each set of criteria emphasizes a categorical approach to diagnosis, despite patients who often show more than one problem—typically AD and cerebrovascular problems¹². While each set of criteria allows for “mixed diagnoses”, they appear to underestimate the role of cerebrovascular lesions in both the risk of AD and the degree of its expression¹³.

In practice, each set of criteria sees AD chiefly in cognitive terms. Another way to think of dementia, however, is as a decline of “effective behaviour”¹⁴—a phenomenon seen often in clinical practice, in which cognitive and non-cognitive changes can compete for prominence as the complaints of patients and families. Perhaps it is this experience which underlies the recognition of non-cognitive features in the ICD-10 clinical description, or in the DSM-IV subtypes. Nevertheless, these non-cognitive aspects have not much informed our research understanding of disease presentation, and neither are they likely to. Given the ongoing search for clinical correlates of structural or chemical preclinical features, cognitive deficits (which are readily quantifiable) are likely to achieve even

greater prominence. Both the near-continuous distribution in neuropsychological test performance scores across an unimpaired to demented range^{15,16} and the preference for categories mean that we are more likely to see a renewed “cut points” debate than a debate over the meaning of cognitive vs. non-cognitive symptoms as the basis for future revisions of diagnostic criteria for AD.

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The Neuropathology of Alzheimer's Disease

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Alzheimer's disease (AD) is the most common cause of dementia at any time of life and, although the brains of patients with this disorder show remarkable pathological changes, the nosological status of these and their relevance to the underlying neurodegenerative process has been the subject of vigorous debate. In this chapter the gross and microscopical changes of AD will be described.

IMAGING STUDIES

Structural imaging, by CT or MRI, fails to reveal changes in the brain specific to AD. Usually there is evidence of widespread cortical atrophy and ventricular enlargement, although not in a way that obviously discriminates AD from other neurodegenerative dementing disorders. A more severe medial temporal atrophy has been claimed to distinguish AD from normal ageing¹, although this does not differentiate between AD and other disorders, such as frontotemporal dementia, where a similar degree of hippocampal atrophy is seen^{2,3}. Functional imaging, using PET or SPECT, reveals a conspicuous biparietal deficit in many patients with AD, although not all individuals show this and this pattern can also be seen in patients with Lewy body dementia⁴. Hence, definitive diagnosis still remains the province of histopathology, although this is usually made retrospectively following the death of the affected person.

THE AUTOPSY APPEARANCE OF THE BRAIN

Although the pattern of atrophy in AD may be quite distinct from that of the normal elderly^{2,5} and of other dementing disorders of diverse aetiology^{3,6} the diagnosis cannot be made at autopsy from visual inspection alone. Nonetheless, brain weight is usually reduced due to cortical atrophy (shrinkage of the gyri and widening of the sulci), which can be widespread but most often is severe in the medial temporal regions, particularly the parahippocampal gyrus, while the occipital lobe and the motor cortex are generally spared².

The cortical grey matter is reduced in thickness and the white matter, while macroscopically normal in appearance, is lost proportionately to that of the grey matter². The ventricular system is dilated, most markedly in the temporal horns of the lateral ventricles. The substantia nigra usually shows normal pigmentation but frequently there is loss of pigment from the locus caeruleus. Cerebrovascular changes are often coincidentally present, but do not necessarily indicate a multi-infarct dementia.

HISTOPATHOLOGICAL CHANGES

The histopathology of AD displays several abnormalities and, although these changes can also be found in the brains of most normal elderly individuals, it is their greater extent and severity and regional pattern of distribution that is characteristic of AD.

Amyloid Plaques

These are aggregates (plaques) of an amyloid protein—amyloid β protein ($A\beta$)—typically 50–200 μm in diameter, within the cerebral cortex and other grey matter regions of the brain. The $A\beta$ may be surrounded by abnormal presynaptic nerve cell processes called dystrophic neurites. The same protein is often present within the walls of leptomeningeal and intracortical blood vessels, causing an amyloid angiopathy. Four plaque types have been identified⁷:

1. *Diffuse* plaques have even $A\beta$ deposition with ill-defined borders, not forming discrete rounded masses. They are the most common plaque type in AD and can only be defined by immunohistochemical staining with antibodies to $A\beta$; they have no dystrophic neurites.
2. *Primitive* plaques are discrete rounded $A\beta$ deposits without a dense core but having some dystrophic neurites. They can be stained using conventional amyloid stains.
3. *Classical* plaques have a dense star-shaped $A\beta$ core surrounded by a corona of radiating wisps of $A\beta$ and many dystrophic neurites.
4. *Compact* or *burnt-out* plaques have dense $A\beta$ cores without a surrounding corona, or neurites.

Primitive and classical plaques with dystrophic neurites containing PHF are termed “neuritic plaques”, synonymous with what used to be known as “senile plaques”. Synapse loss occurs from primitive and classical plaques, but not diffuse plaques^{6,2}.

Neuritic, but not diffuse, plaques contain reactive astrocytes and activated microglial cells^{8,9}; the latter may play a role in the “processing” of $A\beta$ during plaque evolution. Although unproved, the prevailing view is that amyloid plaques undergo, during their prolonged life history, a series of evolutionary changes from diffuse to cored plaques. This involves not only compositional changes in $A\beta$, but also changes in associated glial cells. This evolutionary process has largely been deduced from studies of persons with trisomy 21 (Down's syndrome) dying at different ages^{9,10}, who inevitably develop the histopathology of AD if they live past 50 years of age.

Although plaques are always most dense in cerebral cortex, they can also be numerous in white matter and subcortical areas, including the basal ganglia and cerebellum, where the deposits may be associated with dystrophic neurites but not PHF¹¹. Plaques are common in the cerebral cortex in normal ageing, even in high density in some cognitively normal elderly patients^{12,13}, but are generally of the diffuse or primitive types^{14,15}. The number of plaques, especially those containing $A\beta_{40}$, is increased in a gene dose-dependent way in the presence of the $\epsilon 4$ allele of the apolipoprotein gene^{16,17}.

$A\beta$ is derived, by proteolytic cleavage, from a large precursor, the amyloid precursor protein (APP). This is a transmembrane protein with a large extracellular amino-terminal portion and a small intracellular carboxy-terminal stub. $A\beta$ protein is present in plaque amyloid as a heterogeneous mix of cleaved APP fragments, 39–43 amino acids long^{18,19}.

APP is catabolized by enzymes known as “secretases”. One enzyme, α -secretase (a metalloprotease)^{20,21} cleaves APP across the middle of its transmembrane domain and in this way $A\beta$ formation is not possible. Another enzyme, β -secretase (a transmembrane aspartic protease), termed β -site APP-cleaving enzyme (BACE)²², cuts APP at the amino-terminus of the $A\beta$ domain. The $A\beta$ peptide can then be released from this carboxy-terminal stub by a third enzyme, γ -secretase—which requires presenilin-1 protein for its activity, acting around amino acids 40–43 of the $A\beta$ sequence. The major catabolic product of these enzyme activities is $A\beta_{40}$, with lesser quantities of the longer peptide $A\beta_{42}$. However, because of a higher propensity to form amyloid fibrils the predominant, and indeed the sole, $A\beta$ species in diffuse plaques of the cerebral cortex, corpus striatum and cerebellum is $A\beta_{42}$: $A\beta_{40}$ is mostly present within the cored, neuritic plaques²³. Paradoxically, the predominant peptide within blood vessel walls is $A\beta_{40}$, with variable, and usually much lesser, amounts of $A\beta_{42}$ ²⁴. Studies on Down’s syndrome²⁵ indicate that $A\beta_{42}$ is the earliest peptide deposited, $A\beta_{40}$ deposition occurring subsequently within that subset of diffuse plaques evolving into cored plaques²³.

Some cases of familial AD result from point mutations in the APP gene (APP_{670/671} and APP₇₁₇) on chromosome 21²⁶, whereas in others mutations in genes on chromosomes 14 and 1, known as presenilin-1 and presenilin-2, respectively, are responsible²⁷. Clinicopathological studies indicate that these familial AD cases have an especially high tissue deposition of $A\beta$, particularly $A\beta_{42}$, and very severe amyloid angiopathy^{28,29}. Facilitation of $A\beta$ deposition, particularly $A\beta_{42}$, may be the mechanism whereby the APP and presenilin mutations operate. A different mutation in the APP gene (APP₆₉₃) causes hereditary cerebral haemorrhage with amyloidosis (Dutch type)³⁰.

Neurofibrillary Tangles (NFT)

NFT are abnormal filamentous inclusions that form inside nerve cells. They are not specific to AD and are seen in other neurodegenerative disorders, as well as being present (in lower numbers) in the brains of elderly non-demented persons, particularly in the hippocampus and temporal cortex^{31,32}. Typically, NFT are found in temporal, frontal and parietal cortical areas with sparing of the paracentral (sensory and motor) and occipital cortices. They occur in subcortical areas such as the hypothalamus and nucleus basalis of Meynert, as well as in the dorsal raphe and locus coeruleus. When NFT formation in these regions is severe, NFT can be seen in the substantia nigra, thalamus and basal ganglia¹⁸.

Traditionally, NFT are demonstrated by silver staining, but they can also be detected immunohistochemically using antibodies to their constituent proteins. The classical NFT is a flame-shaped skein of fine fibrils occupying much of the cell body. In haematoxylin and

eosin-stained preparations, NFT are faint basophilic fibrillar inclusions. Certain NFT, termed “ghost tangles”, appear eosinophilic and represent liberated neuropil remains following death of the affected neurone. Rounded ball-like NFT are termed “globose tangles” and are most common in neurones of the brainstem nuclei, but also in smaller cortical neurones¹⁸.

Ultrastructurally, NFT are composed of pairs of filaments, diameter 20 nm, twisted with a periodicity of 80 nm, hence termed paired helical filaments (PHF)³³. Evidence suggests them to be a ribbon-like double helical stack of transversely-arranged subunits³⁴.

NFT are composed of at least two major proteins, tau^{35–37} and ubiquitin^{38,39}. Tau protein stabilizes the microtubular cytoskeleton. The tau protein in PHF is abnormally phosphorylated^{40,41} and abnormalities in protein phosphorylation may be fundamental to the neurodegenerative process of AD⁴². Ubiquitin labels cell proteins for degradation⁷⁰ and its presence in PHF is presumably targeting the abnormal protein for (attempted) degradation.

Like plaques, NFT undergo evolutionary changes that correlate with their morphology and immunoreactivity⁴³:

Stage 0: amorphous tau immunoreactivity without aggregation into filaments (pre-tangles).

Stage 1: delicate fibrillar structures immunoreactive for tau and ubiquitin.

Stage 2: skeins and whorls of densely aggregated filaments, immunoreactive for tau and (sometimes) ubiquitin, filling the nerve cell body and displacing the nucleus.

Stage 3: extracellular NFT without tau immunoreactivity but ubiquitin-positive. The NFT is infiltrated by astrocytic processes, and $A\beta$ may be deposited upon the filaments⁴⁴.

The progress of AD can be staged by following the spread of NFT from hippocampal regions to neocortex and subcortex⁴⁵.

Neuropil threads are abnormal nerve cell processes (dystrophic neurites) in the neuropil of the cerebral cortex, not associated with plaques⁴⁶. They are swollen, distorted structures containing PHF⁴⁷, antigenically similar (for tau and ubiquitin) to, and probably contiguous with, the NFT in neurones. These abnormal processes may interfere with neuronal communication, contributing to the cortical deficit.

The importance of neurofibrillary pathology in AD has been underscored by the discovery that similar abnormal tau aggregates occur in neurones and glia in frontotemporal dementia⁴⁸, due to mutations in the tau gene^{49–51}. These mutations alter the microtubule binding capacity of tau, favouring self-assembly of the mutated protein into pathological structures, or they change the expression pattern of the tau gene, leading to intracellular accumulations of tau isoforms with four microtubule binding repeat domains. Pathological tau aggregates are also formed in a related condition, progressive supranuclear palsy⁵², in the absence of $A\beta$ deposition. Neurodegenerative disease leading to dementia can therefore be caused by neurofilamentous aggregations of tau *alone*, irrespective of whether amyloid plaques are present. That is not to say, however, that $A\beta$ deposition in AD is of no importance. This may represent the route towards the formation of abnormal tau in this particular disorder, whereas in other conditions other pathways to tau pathology are invoked. Nonetheless, once pathological tau is formed and accumulated, a common mechanism of cell death may be set up in all “tauopathies” and this clearly has profound therapeutic implications for AD and other dementing disorders.

Granulovacuolar Degeneration (GVD)

GVD is an accumulation of membrane-bound vesicles, containing amorphous material, within nerve cells. In haematoxylin and

eosin- or silver-stained sections GVD appears as numerous dot-like particles, each with a surrounding clear halo. These bodies may also be seen in ageing brains, but their number and tissue density are considerably greater in AD. Some granulovacuoles are immunoreactive for ubiquitin^{53,54} and they may represent residual bodies from lysosomal-mediated proteolysis⁵⁵.

Hirano Bodies

These are rod-shaped bodies, 15 μm wide and 60–100 μm long, that appear in neurones with ageing but, again, with greater density in AD. They are brightly eosinophilic and immunohistochemistry suggests they are derived from the cytoskeletal protein, actin⁵⁶.

Neuronal loss in AD

There is conspicuous loss of neurones from the cerebral cortex and hippocampus in AD, particularly large pyramidal cells^{57–59}. This loss is more marked in younger patients (<80 years), although it is still significant in older individuals⁵⁹. Golgi studies show loss of dendritic arborization in surviving neocortical cells⁶⁰ and electron microscopy and immunohistochemistry indicate considerable synapse loss^{61–63}. Cell loss from the nucleus basalis of Meynert, the cholinergic input into the cortex, is reflected neurochemically by reduced choline acetyltransferase, and that from the locus coeruleus can be related to a decline in cortical noradrenaline⁶⁴.

Amyloid Angiopathy

The deposition of A β in cerebral arteries, termed “amyloid angiopathy” or “congophilic angiopathy”, is almost invariable in AD but can also exist in its own right. This amyloid angiopathy may sometimes be responsible for (lobar) cerebral haemorrhage, a cause of secondary stroke in AD^{65–67}.

White Matter Changes

A reduction in the amount of white matter in AD is associated with a decrease in the intensity of myelin staining. This has been described as incomplete infarction^{68,69} and, although it may mimic Binswanger’s disease pathologically and on imaging, it is not associated with lacunar infarcts or hypertensive arteriosclerosis. Such white matter changes may sometimes be related to ischaemia due to amyloid angiopathy, but bouts of systemic hypotension might also be causal.

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Oxford Project to Investigate Memory and Ageing (OPTIMA): a Longitudinal Clinicopathological Study of Dementia and Normal Ageing

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OPTIMA was founded in 1988 by A.D. Smith, K.A. Jobst, E.M.-F. King and M.M. Esiri, with the aim of studying in parallel a cohort of patients with memory problems and age-matched controls. The total number of subjects at the end of 1999 was 666, of which 361 are patients with dementia. OPTIMA is a longitudinal clinicopathological study and its main strength lies in the very high necropsy rate (94% of the 207 who have died), which permits correlation of findings in life with those of neuropathology. Each year each subject has a full clinical examination, lumbar puncture, neuropsychology (the CAMDEX, supplemented by other tests), CT scans (axial and temporal lobe-orientated) and SPECT scans (Ceretek, for cerebral blood flow). A subset of 155 subjects have had annual volumetric MRI scans and more detailed neuropsychology.

The main findings so far are:

- Comparison of standard clinical diagnostic procedures with histopathological diagnosis, showing the poor accuracy of current clinical diagnostic procedures¹.
- Development of a more accurate diagnostic procedure for Alzheimer's disease (AD) in life by a combination of structural (CT) and functional (SPECT) brain imaging².
- Recognition that AD is distinct from ageing and is a true disease that follows a "catastrophic event" in the brain, leading to atrophy of the medial temporal lobe³.
- Discovery of a biological "state" marker, the thickness of the medial temporal lobe, that can be used to follow the progression of AD⁴.
- Discovery that nerve cells in AD brain express markers of the cell division cycle⁵.
- Recognition of the additive effect of minor cerebrovascular disease and AD-type pathology in clinical dementia^{6,7}.
- Identification of a gene (K variant of butyrylcholinesterase) that markedly increases the risk of AD in those who also have the ApoE4 gene⁸.
- Discovery of a risk factor (elevated blood levels of homocysteine) for AD and for vascular dementia that is potentially modifiable by diet⁹.
- Finding that raised blood homocysteine levels are associated with low performance on cognitive tests in the normal elderly¹⁰.

Projects currently under way include the search for further genetic and non-genetic risk factors for dementia, with particular emphasis on modifiable risk factors; a pilot clinical trial of high-dose folic acid and vitamin B₁₂ in subjects with dementia (together

with Professor G. Wilcock, Bristol); development of novel memory tests to detect pre-symptomatic AD; use of sub-voxel co-registration MRI scans to follow progression of AD over a period of a few months (with Professor G. Bydder, Hammersmith Hospital).

The current Director of OPTIMA is Professor A. David Smith, the Clinical Director is Professor Robin Jacoby and the Operational Manager and Senior Nurse is Mrs Elizabeth King. Funding is from Bristol-Myers Squibb, the Medical Research Council, National Health Service R&D and several charities.

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Consortium to Establish a Registry for Alzheimer's Disease (CERAD)

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CERAD was funded by the National Institute on Aging in 1986 to develop a battery of standardized instruments for the evaluation of patients with Alzheimer's disease (AD). Until that time clinical investigation of AD and comparison of research findings was hampered by the absence of standardized assessment and uniform diagnostic criteria. The assessments developed by CERAD have been evaluated on over 1000 patients with AD and nearly 500 control subjects, seen at 24 major University medical centers across the USA. Because of their sensitivity to dementia, they have also been used in epidemiologic surveys of the elderly, to aid in identification and staging of those with dementia. The measures permit uniform identification of dementia and standardized assessment of AD.

CERAD has developed and evaluated three primary assessments. These include a clinical battery, a neuropsychological battery, and a neuropathological assessment. An overview of the contents of these assessments is given in Table 1. In addition, specialized assessments have been developed to assess family history of AD, Parkinson's disease, and Down's syndrome; extrapyramidal dysfunction in AD; neuroimaging; behavioral pathology; and assessment of service use.

Videotapes demonstrating administration of these measures are available. Educational brochures on memory loss, AD, the importance of autopsy, and an autopsy resources packet to help sites, have been prepared. Many of the CERAD assessments have been translated into various European and Asian languages, and are in use internationally.

A brief but extensive review of CERAD, including a bibliography covering the first 10 years, has been published¹. Multiyear data on CERAD patients and control subjects as well as the CERAD measures, are available on CD-ROM. This and additional information can be obtained by writing to the Principal Investigator, Dr A. Heyman, CERAD, Box 3203, Duke University Medical Center, Durham, NC 27710, USA.

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Table 1. Overview of the contents of the primary CERAD assessments

<i>Clinical battery</i>	<i>Neuropsychological battery</i>
Demographic data on subject and informant	Verbal fluency
Clinical history, including cognitive function, systemic disorders, cerebrovascular history, parkinsonism, depression, drug effects	Modified Boston Naming Test
Blessed Dementia Scale (ADL)	Mini-Mental State Examination
Screen for Behavior Rating Scale for Dementia (BRSD)	Word list memory
Short Blessed, Calculation, Clock, Language	Constructional praxis
Clinical examinations, including brief physical, overall neurological assessment, extrapyramidal dysfunction	Word list recall
Laboratory and imaging studies	Word list recognition
Clinical diagnosis, including CDR staging, diagnostic impression for:	Constructional praxis recall
Possible dementia prodrome	The following are used as needed:
Probable and possible AD	Shipley Scale
Non-AD dementias	Wechsler Memory Scale, Paired Association I
	Trail Making, A and B
	Wechsler Memory Scale, Paired Association II
	Nelson Adult Reading Test
	Finger tapping
	Verbal fluency (F and P words)
	<i>Neuropathology Assessment</i>
	Demographic information and history
	Gross examination
	Cerebrovascular disease (gross)
	Microscopic findings: vascular; major non-vascular; hippocampus and neocortex
	Neurohistologic findings
	Neuropathological diagnoses
	Final assessment

Neurotransmitter Changes in Alzheimer's Disease: Relationships to Symptoms and Neuropathology

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Cell death and histopathological changes affecting a number of neuronal systems are considered to result in the development of the typical symptomology of Alzheimer's disease (AD), characterized by gross and progressive impairments of cognitive function, which are often accompanied by behavioural disturbances such as aggression, depression, psychosis, apathy and wandering. Such non-cognitive behavioural symptoms are also considered to relate to structural and functional alterations in neurotransmission. Carers find behavioural disturbances difficult to cope with and the presence of such behaviours in AD patients often leads to the need for institutionalization¹. The challenge has been to identify changes in specific neurotransmitter systems that underlie cognitive impairment and particular behavioural problems and to develop rational therapeutic strategies.

NEUROCHEMICAL AND HISTOPATHOLOGICAL CHANGES IN AD

The majority of biochemical studies of AD have relied on information derived from post mortem brain which typically represents the late stage of the disease (8–10 years after onset of symptoms). In these studies there is considerable evidence of gross brain atrophy, histopathological features and multiple neurotransmitter abnormalities affecting many brain regions. However, investigations of biopsy tissue taken from AD patients 3–5 years (on average) after the onset of symptoms indicate that a selective neurotransmitter pathology occurs early in the course of the disease².

Acetylcholine

Changes affecting many aspects of the cholinergic system in patients with AD have been reported since the initial discovery of deficits in choline acetyltransferase activity in post mortem brains^{3–5}. In biopsy samples from AD patients, presynaptic markers of the cholinergic system were also uniformly reduced². Thus, choline acetyltransferase activity, choline uptake and acetylcholine synthesis are all reduced to 30–60% of control values. The clinical correlate of this cholinergic deficit in AD was, until recently, considered to be cognitive dysfunction. Such a conclusion was supported by clinicopathological studies in AD and parallel experiments in non-human primates or rodents, which demonstrated disruptive effects of basal forebrain

cholinergic lesions on cognitive functions. Such studies led to the "cholinergic hypothesis of geriatric memory dysfunction"⁶.

Furthermore, cholinergic deficits in AD occur to the greatest extent in cortical areas primarily concerned with memory and cognition—the hippocampus, adjacent temporal lobe regions and select frontal areas. In a recent study⁷ regional variations in the loss of cholinergic fibres in AD were assessed on the basis of acetylcholinesterase (AChE) histochemistry. Greatest fibre loss (> 75%) was apparent in temporal association areas, with various frontal areas, including granular orbitofrontal, dysgranular orbitofrontal, prefrontal association, frontal operculum, prefrontal association and frontal pole, demonstrating fibre losses in the range 45–75%. In other cortical areas, including primary motor, premotor association, anterior and posterior cingulate, fibre loss was less than 45%.

Neuropathologically, loss of neurons from the nucleus of Meynert (Ch4 cholinergic nucleus) is well documented in AD, although the extent of the loss reported varies from moderate to severe, and it has been suggested that in AD cholinergic dysfunction exceeds degeneration⁸. Detailed analysis of subpopulations of cholinergic perikarya in the nucleus basalis have been reported by Mesulam and Geula⁹, who identified selective cell loss in Ch4p (the posterior section projecting to temporal cortex). In the intermediate sector, Ch4id, which includes projections to the frontal cortex, neuron loss is not as extensive, consistent with the moderate loss of cholinergic enzyme activity.

On the basis of the above evidence, neocortical cholinergic innervation appears to be lost at an early stage of the disease and this is supported by a recent study¹⁰ in which the cholinergic deficit (reduced ChAT activity) has been related to Braak staging. Braak stages I and II are considered to represent the earliest presentation of AD, with neurofibrillary tangles in the entorhinal cortex, and a 20–30% loss of ChAT activity was reported in brains from patients at these stages of AD¹¹. However, another study using the Clinical Dementia Rating Scale (CDR) suggests that the greatest reduction in markers of the cholinergic system occurs between moderate (CDR 2.0) and severe (CDR 5.0) disease, with little change between non-demented and the mild stage (CDR 0–2)¹².

There has been a recent shift of emphasis regarding the clinical significance of cholinergic deficits. Non-cognitive or neuropsychiatric, in addition to cognitive, symptoms also appear to have a cholinergic component¹³. For example, visual hallucinations relate to neocortical cholinergic deficits¹⁴, such deficits (e.g. loss of ChAT) being greater in Lewy body dementia (DLB), where

hallucinations are common, than in AD where they are less common¹⁵. Reductions in cortical ChAT activity in patients with dementia, in addition to correlating with cognitive decline, are also related to overactivity and aggressive behaviour¹⁶.

It has also been suggested that acetylcholine is centrally involved in the process of conscious awareness¹⁷, and that the variety of clinical symptoms associated with cholinergic dysfunction in AD and related disorders reflects disturbances in the conscious processing of information. There is evidence that implicit memory, for example (which does not involve conscious awareness), is relatively intact in AD^{18,19}.

Glutamate

Loss of synapses and pyramidal cell perikarya (both considered to be markers of glutamatergic neurones) from the neocortex of AD patients correlate with measures of cognitive decline². Although neurochemical studies of glutamate neurotransmission have failed to demonstrate extensive alterations, this may be related to the difficulty in distinguishing the transmitter pool of glutamate from the metabolic pool. Nevertheless, glutamate concentration was reduced by 14% in temporal lobe biopsy samples and by 86% in the terminal zone of the perforant pathway at autopsy of AD patients²⁰. Uptake of D-aspartate, a putative marker of glutamatergic nerve endings, is also reduced in many cortical areas in AD brain². Thus, additional factors other than impaired cholinergic function are likely to contribute to cognitive impairment in AD. However, it is important to remember that glutamatergic neurones of the neocortex and hippocampus are influenced by acetylcholine through nicotinic and muscarinic receptors². Thus, treatment of patients with cholinomimetics is likely to increase glutamatergic function.

Other neurotransmitters

Using biopsy samples from AD patients, serotonergic and some noradrenergic markers are affected, whereas markers for dopamine, γ -aminobutyric acid (GABA) or somatostatin are not altered. When post mortem studies of AD brain are considered, many neurotransmitter systems, including GABA and somatostatin, are involved or are affected to a greater extent². Based on post mortem studies, however, changes in serotonergic neurotransmission may be linked to the behavioural disturbances of AD, such as depression, rather than cognitive dysfunction. For example, patients with AD who were also depressed had lower numbers of serotonin reuptake sites in the neocortex than did patients without this symptom²¹. Furthermore, both reduced serotonergic^{22,23} and increased noradrenergic activities and sensitivity^{24,25} have been linked to aggressive behaviour.

Neurotransmitter receptors

Many neurotransmitter receptors appear to be unaltered in AD; however, studies have demonstrated a reduction in the number of nicotinic and muscarinic (M2) ACh receptors, most of which are considered to be located on presynaptic cholinergic terminals. Despite continuing, often unconfirmed, reports of changes in one or more of the muscarinic receptor subtypes (M₁–M₅), it is generally agreed, on the basis of autopsy studies, that the M1 subtype is unchanged until later in the disease when it may decline, probably in relation to cholinergic (postsynaptic) neurodegeneration. The status of the other subtypes is not clearly established, primarily due to the lack of specific pharmacological labels. Results using antibodies against the different receptor

subtypes, while specific, are complicated by discrepancies between the distribution and density of immunoreactive proteins and localized functional receptors. With respect to muscarinic receptor coupling to G-proteins, most studies using a variety of investigative procedures have identified some degree of uncoupling, especially with respect to the M1 receptor²⁶.

A highly consistent receptor abnormality in AD is the loss of the nicotinic receptor^{27,28}, which appears to primarily reflect loss of the α 4-containing subtype (generally associated with β 2) as opposed to α 3 or α 7 subtypes²⁹. Immunohistochemically, loss of α 4 and β 2 reactive fibres has been observed in temporal cortex, associated with reactive neuropil threads, tangles and plaques³⁰.

NEUROIMAGING

With respect to the cholinergic deficit, whilst measurements of CSF acetylcholine, choline and acetylcholinesterase have been reported in AD, such reports are either unconfirmed or inconsistent. More promising and potentially diagnostic findings have been obtained using *in vivo* functional imaging. The vesicular acetylcholine transporter and acetylcholinesterase, imaged using PET and iodobenzovesamicol and *N*-methylpiperidin-4-yl propionate, respectively, are both reduced in AD patients, and furthermore relate to reductions in MMSE^{31,32}. Using SPECT, muscarinic QNB binding is reduced in advanced but not early cases³³, and iododexetamide, with preference for M2, is also reduced in mild/early cases³⁴. Reductions in nicotine binding have also been detected using PET³⁵. *In vivo* observations relating to non-cholinergic systems (principally noradrenaline, 5-HT and dopamine) have been equally inconsistent regarding CSF parameters and, with respect to neuroimaging markers, only dopaminergic markers have so far been investigated and there is, as expected, no consistent abnormality of the transporter or D2 receptor.

LINKS BETWEEN NEUROTRANSMISSION AND NEUROPATHOLOGY

Mismetabolism of amyloid precursor protein (APP) leading to increased production of β -amyloid has been proposed as the critical event in both familial and sporadic AD causing other changes (tangles, neurone loss, synapse loss and neurotransmission dysfunction). Cholinergic neurotransmission may be a specific target for β -amyloid, since it has been shown to reduce both choline uptake and acetylcholine release *in vitro*³⁶. Furthermore, β -amyloid is reported to bind with high affinity to the α 7 subtype of the nicotinic receptor, suggesting that cholinergic function through this receptor may be compromised because of high levels of (soluble) peptide in AD brains³⁷.

There is increasing evidence that various neurotransmitter systems are capable of influencing the metabolism of APP, favouring the non-amyloidogenic processing³⁸. In particular, stimulation of muscarinic M1 receptors increases APP secretion, while decreasing β -amyloid production³⁹. Furthermore, nicotinic receptor stimulation is associated with reduced plaque densities in human brain⁴⁰. These results suggest that compounds being developed for symptomatic treatment may have a serendipitous effect on the continuing emergence of pathology by reducing the production of β -amyloid.

CHOLINERGIC APPROACHES TO TREATMENT

Biochemical studies of postmortem brains from AD patients showing evidence of a substantial presynaptic cholinergic deficit,

which correlated with cognitive impairment⁴¹, together with the emerging role of ACh in learning and memory⁶, clearly suggested a rational approach to treatment. However, more recent studies have identified a role for the cholinergic system in attentional processing rather than memory.

A prediction of the cholinergic hypothesis is that drugs that potentiate central cholinergic function should improve cognition in AD patients. There are a number of approaches to the treatment of the cholinergic deficit; however, the use of acetylcholinesterase inhibitors is the most well-developed approach to the treatment of AD to date⁴².

During the late 1980s and early 1990s, the first cholinomimetic compound, tacrine, underwent large-scale clinical studies and established clearly the benefits of ChE treatment in patients with a diagnosis of probable AD. Tacrine was subsequently approved for use in some, but not all, countries. Statistically significant, dose-related improvements on objective performance-based tests of cognition, clinician- and caregiver-rated global evaluations of patient well-being and also quality of life measures have been reported⁴³. Unfortunately, potentially serious adverse side effects have limited the use of this compound.

A so-called second generation of ChE inhibitors has been developed, including donepezil, rivastigmine, metrifonate and galantamine⁴². Such compounds demonstrate a clinical effect and magnitude of benefit of at least that reported for tacrine, but with a more favourable clinical profile. Furthermore, evidence is emerging from clinical trials of cholinomimetics that such drugs may improve the abnormal non-cognitive, behavioural symptoms of AD. The cholinesterase inhibitors physostigmine, tacrine, rivastigmine and metrifonate have variously been reported in placebo-controlled trials to decrease psychoses (hallucinations and delusions), agitation, apathy, anxiety, disinhibition, pacing and aberrant motor behaviour and lack of cooperation in AD^{13,44}. In a recent open-label trial of Exelon (rivastigmine), patients with DLB almost all responded positively on one or more of these measures⁴⁵.

CONCLUSIONS

In AD many different pathological manifestations, such as cortical and subcortical β -amyloidosis (plaques), abnormal tau (tangles and dystrophic neurites), neuronal and synapse loss and various transmitter deficits, provide an increasingly complex framework for clinical-neuropathological correlations. It is therefore unlikely that cholinergic deficits alone will account for the full spectrum of cognitive and non-cognitive symptoms seen in AD. In these circumstances it is perhaps surprising that cholinomimetic therapy has been a modest success in many patients, improving cognitive and non-cognitive symptoms and activities of daily living. Such therapy is all that is available at present and may well still be required as new therapies designed to slow disease progression come into the clinic.

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Antemortem Markers

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INTRODUCTION

It has become increasingly clear that the disease process of Alzheimer's disease (AD) is multifaceted. It is thus difficult to single out any particular factor as the root cause of the disease, since AD appears to be a complex disorder involving several genes interacting with environmental factors. Apart from the familial forms of the disease, in which the gene mutations have been elucidated, many proteins, enzymes and other factors are involved in the process of neurodegeneration, in the formation of plaques and tangles and in the development of an inflammatory state of the brain. In all, more than 100 proteins and other factors have been found to be altered in AD patients compared with controls.

For all that, very few markers suitable for antemortem diagnostic purposes have emerged, since many of the above-mentioned alterations are not specific for AD, while others pertain only to subsets of AD¹. While it has been argued that the most powerful antemortem marker in AD is a clinical diagnosis based on an adequate range of observations², such diagnosis is at present to some degree still one of exclusion. An ideal biological marker would allow for greater specificity and sensitivity than clinical diagnosis, and be readily obtainable. While neuropathological biopsy diagnosis of AD allows specificity and sensitivity, it is rarely clinically warranted or available. The neurobiological alterations present in AD may be reflected in changes in cerebrospinal fluid (CSF) neurotransmitters or neurochemicals, or in a change in systemic tissues, including blood constituents. It should be borne in mind, however, that CSF measurements are influenced by a variety of factors, including CSF gradients, age and sex, diurnal and seasonal variation, state of the blood-brain barrier, blood contamination, contributions from the spinal cord, phase of illness, psychomotor activity, stress and diet. Measurement of blood constituents may also reflect concentration differences due to diurnal rhythms and other factors.

In this chapter on antemortem markers, we will briefly review neurotransmitters and neurochemistry, systemic pathology and brain imaging.

NEUROTRANSMITTERS AND NEUROCHEMISTRY

The Cholinergic System

The most effective drugs so far for the treatment of AD are the acetylcholinesterase (AChE) inhibitors. The introduction of these agents followed the discovery that cholinergic neurons were depleted and that cholinergic function was significantly decreased in the basal forebrain of AD patients³. CSF markers of

cholinergic function have been studied, but have not yielded consistent results. For example, measurements of AChE and pseudocholinesterase (PChE) have led to the conclusion that cholinergic basal forebrain neurons are not a major source of cholinesterases in the CSF and do not provide evidence for using CSF cholinesterases as a diagnostic marker of basal forebrain cholinergic cell loss⁴.

The Noradrenergic System

Autopsy studies of AD brains demonstrate loss of cells in the locus coeruleus, the major nucleus of origin of noradrenergic fibres. Reduced noradrenaline (norepinephrine NE) in autopsy samples of AD brains has been a fairly consistent finding. In contrast, CSF and plasma NE and 3-methoxy-4-hydroxyphenylethylene glycol (MHPG) appear significantly higher in patients with advanced AD than in patients with moderate AD or controls^{5,6}. Patients with advanced AD have not only biochemical indices of noradrenergic hyperactivity but also physiological pointers to this, including higher heart rate and blood pressure. AD patients with the most severe dementia have the greatest rise in CSF MHPG levels following administration of probenecid. There is also evidence for blunted growth hormone response to clonidine in AD patients, suggestive of altered α -2-adrenergic receptor sensitivity⁷. It may be hypothesized that increased activity and turnover of the noradrenergic system may compensate for cell loss and that a limited number of NE cells remain highly active in AD patients. Severe neuronal loss in advanced AD may lead to a compensatory increase in locus coeruleus firing rate, contributing to symptoms such as pacing, agitation, insomnia and weight loss.

The Serotonergic System

Numerous autopsy studies of AD brains have suggested a serotonergic deficit. Although there have been reports that the major serotonin metabolite, 5-hydroxyindoleacetic acid (5-HIAA), is unchanged in the CSF of AD patients, most studies indicate a reduction in CSF 5-HIAA. In one study that demonstrated significantly lower mean 5-HIAA levels in AD, the wide variability in values suggested that the changes were non-specific, secondary to the cerebral degeneration in AD⁸. There is also evidence of increased behavioural sensitivity to *m*-chlorophenylpiperazine in AD patients, consistent with damage to serotonin pathways⁹.

Melatonin, the pineal hormone biosynthesized from serotonin, has been demonstrated to be significantly decreased in the CSF of elderly patients, and more so in AD patients. In elderly (> 80 years

of age) non-demented subjects, CSF melatonin levels were half those of younger (age 41–80) control subjects, and in AD patients the CSF melatonin levels were only one-fifth of those found in control subjects ($p < 0.0001$)¹⁰.

Other neurochemical systems have been investigated as potential antemortem markers¹¹. Somatostatin is reduced in AD brains and CSF somatostatin levels have been found to be significantly lower in patients with AD compared to controls. These findings do not, however, appear specific to AD. Other CSF peptidergic findings in AD include decreased vasopressin, decreased thyrotropin-releasing hormone, and decreased delta sleep-inducing peptide. Another system that may be disordered in AD brains comprises trophic factors such as gangliosides.

SYSTEMIC PATHOLOGY

It is possible to view AD as a systemic illness. Thus, if AD were a genetic disorder, then disturbances at the molecular level may be expressed in non-neural tissue, with systemic effects.

Studies of blood cell cholinergic function have been undertaken since central cholinergic dysfunction in AD became a favoured hypothesis. Studies of red blood cell (RBC) and plasma choline and AChE do not support their validity as antemortem markers. However, studies indicate differences in the dynamics of RBC choline uptake, suggesting a vulnerability of cholinergic neurons in patients with AD¹².

Fluidity of the platelet membrane (PMF) is increased in patients with AD compared to patients with vascular dementia (VaD) and elderly controls^{1,13,14}. Only about 50% of AD patients demonstrate this abnormality; however, increased PMF appears to be a familial trait, and the subgroup of AD patients in whom it manifests suffer from an earlier onset and a more rapidly progressive decline¹⁵. In a prospective longitudinal study to evaluate PMF as a putative risk factor for AD, nine of 330 people with increased PMF (initially asymptomatic first-degree relatives of probands with AD) developed AD after 7.5 years¹⁶. On a biochemical level, it was found that free radical-induced lipid peroxidation increased the fluidity of platelet membranes, providing a postulated mechanism underlying increased PMF¹⁷.

In fibroblasts, abnormalities in enzymatic activity, glucose metabolism, abnormal calcium metabolism, impaired DNA repair and potassium channel dysfunction¹⁸ have been observed.

Inflammation

Neuropathological findings in AD brains show an inflammatory response. CSF interleukin-1 β (IL-1 β) concentrations were significantly higher in AD patients than in patients with VaD, normal pressure hydrocephalus or multiple sclerosis¹⁹. Activated microglia may participate in the initial stages of neurodegeneration. Microglial antibodies in the CSF have been found in AD patients²⁰. CSF microglial antibodies were present in at-risk descendants of familial AD patients, some of whom subsequently developed AD, and in AD patients in contrast with other types of dementia²¹. The presence of microglial antibodies may, however, be non-specific, as a similar mechanism has been implicated in the pathophysiology of VaD. Drugs aimed at diminishing inflammation and the activity of microglia are currently being investigated.

A miscellany of other research areas in AD can conveniently be mentioned in this section. These include studies of visual system dysfunctions, olfactory deficits, extrapyramidal dysfunction, atypical dermatoglyphic patterns, altered sweat response, abnormal glucose tolerance and vitamin B₁₂ deficiency. Such work, although interesting, has not yet led to a sensitive and specific antemortem marker.

Protein Abnormalities

Mutations of the amyloid β -protein precursor (APP) have been found in familial AD with early onset. A prominent feature of the senile plaques found in AD is the deposition of β -amyloid. The other hallmark lesion for AD is the neurofibrillary tangle, which contains tau protein. Given that the clinical phase of AD may be preceded by a 15–30 year period of deposition of amyloid and tau protein, markers to predict the development of AD should ultimately be obtainable²². While the concentration of tau protein in CSF was found to be significantly higher in AD patients compared to non-demented controls, reports of altered APP and β -amyloid CSF concentrations have not been consistent²².

Environmental Factors in AD

Although AD has been linked to a genetic aetiology, studies with monozygotic twins revealed variability in age at onset of as much as 9, 15 or 20 years²³. This led to the suspicion that some environmental factor(s) ingested from food and water, or a deficiency of some protective element, may accelerate disease expression in susceptible individuals. Several metals have been investigated in AD in the hope of finding a marker for the disease. Of these, iron, aluminium, zinc and mercury are considered to be the most important. Aluminium was found in neurofibrillary tangles^{24,25}, aluminium and silicon were found in cores of senile plaques²⁶ and aluminium levels were increased in the hippocampus and cerebral cortex of AD patients compared with non-demented controls²⁷. Aluminium is not easily absorbed by the body, since the gastrointestinal tract forms a major cellular barrier to its absorption. In AD this barrier may be compromised²⁸. The findings of increased aluminium in AD have, however, been inconsistent.

Iron metabolism is altered and the iron transport protein transferrin decreased in the serum of AD patients. Furthermore, a genetic form of transferrin, TfC2, which is associated with diseases attributed to free radical damage, has an increased allele frequency in AD^{29–31}. Serum levels of the iron binding protein p97 (melanotransferrin) were reported to be elevated in AD patients, discriminating between AD patients and controls³², a finding which has to be confirmed. Finally, blood mercury levels are also significantly raised in AD patients³³.

BRAIN IMAGING

There has been considerable interest in the use of brain imaging in the diagnosis of AD (see Section EX). The medial temporal lobe is involved in cognitive functions of the brain and is the region reflecting the most extensive pathological changes in AD. Using temporal-lobe-orientated computed tomography (CT), it was demonstrated that decreased hippocampal width predicted AD with a detection rate of 92% when a cut-off was selected to yield a 5% false-positive rate. These cases were subsequently confirmed histopathologically³⁴. CT scans also revealed that the atrophy of the medial temporal lobe in AD was related to the progression of pathology³⁵.

Magnetic resonance imaging (MRI) studies yielded similar results. On MRI, AD patients and controls were best told apart using left amygdala and entorhinal cortex volumes³⁶. Coupled to blood flow studies using single photon emission computed tomography (SPECT) and using relative left temporoparietal cortex blood flow, the imaging studies yielded 100% discrimination between AD patients and controls³⁶. In addition, in older patients with mild cognitive impairment, hippocampal atrophy determined by premonitory MRI-based volume measurements was

predictive of subsequent conversion to AD³⁷. In distinguishing between AD and other dementias, one study differentiated AD from normal ageing, depression, VaD and other causes of cognitive impairment³⁸, while in another study, no significant volumetric differences were found between patients with subcortical VaD and AD patients, apart from the volume of the cerebellum³⁹.

Positron emission tomography (PET) has been found to differentiate between AD and VaD: in AD the typical metabolic pattern is hypometabolism in temporoparietal and frontal association areas, while in VaD scattered areas of hypometabolism extending over cortical and subcortical structures are seen⁴⁰.

CONCLUSION

It is clear that a number of difficulties beset the researcher interested in finding an antemortem marker for AD. Some of the techniques used in this search have important limitations—in particular, CSF measurements are influenced by a variety of factors. Diagnosis of AD, even at autopsy, may not always be accurate, leading to heterogeneity of the patient sample. In addition, AD may be a heterogeneous disorder, perhaps with a presenile form, but certainly with different manifestations in the much older age group. A marker may be present in a control with the AD trait who has not yet developed AD, and if a marker is related to the severity of AD, it may not distinguish between mild cases and controls. Other dementias may overlap with AD, not only in phenomenology but also in pathogenesis, and therefore display similar markers. There are also mixed cases, where patients, for example, have both AD and VaD.

Nevertheless, the work discussed here does suggest that some combination of measures may ultimately correlate strongly with biopsy diagnosis. Work in molecular and genetic biology and in brain imaging seems particularly promising and may well tie in with our knowledge of neurochemistry and neuropathology. An interdisciplinary effort to find an antemortem marker is useful not only in leading to a better delineation of AD-related dementias and AD subtypes, but also in providing a focus on central pathogenic mechanisms and their possible reversal.

Finally, markers may be expressed differently in different populations. A genetic variant of apolipoprotein E, ApoE ϵ 4 (Chapter 41) has been shown to be a risk factor for late-onset AD. Yet, in spite of relatively high frequencies of ApoE ϵ 4 in African countries, the prevalence of AD is very much lower than in the Western population⁴¹, making it extremely important for us to understand the aetiology of the disease.

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Clinical Features of Senile Dementia and Alzheimer's Disease

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Not all dementias in the senium are forms of Alzheimer's disease (AD), neither does AD always arise in the senium. However, the commonest form of senile dementia and of AD is senile dementia of the Alzheimer type (SDAT). In most countries this alone is thought to account for 50–60% of the senile dementias and, in a form mixed with multi-infarct dementia (MID), for another 15–20%¹.

Alois Alzheimer originally described² a 51 year-old woman whose morbid jealousy was followed by a rapidly progressing amnesia. She displayed paranoid delusions and, it seemed, auditory hallucinations as well as such cognitive defects as disorientation, incomprehension, perseveration, dysphasia, dysgraphia and dyspraxia. She died only 4.5 years after the onset of her disorder, when Alzheimer was enabled to make the neuropathological observations that later caused his name to be given to the disease. Thus "psychiatric" as well as cognitive symptoms were part of the syndrome from the first. Personality change and behavioural disorder are also part of the condition, especially in those referred to psychiatrists.

Most diagnostic criteria for dementia and AD, e.g. the American Psychiatric Association's *Diagnostic and Statistical Manual*, 3rd edn, Revised³ and those of the Royal College of Physicians⁴, require some degree of disability. The operational diagnostic criteria for dementia in the Cambridge Examination for Mental Disorders of the Elderly⁵ include "Progressive failure in performance at work and in the common activities of everyday life—The decline in memory is sufficiently severe to impair functioning in daily life".

Presumably there must be some period of minimal impairment before the disorder becomes disabling, but it is still unclear what are the very earliest features of AD, in its senile or presenile forms. Sufferers tend not to be brought to the notice of the medical or social services until problems arise that jeopardize self-care or strain relationships, when the disease has usually been developing for at least 2 years. Epidemiological surveys using such screening instruments as the Mini-Mental State Examination⁶ have, however, identified some people with mild or borderline cognitive impairment, as have the memory clinics which have been developed in recent years⁷, and some of these have been shown later to have developed dementia. The distinction between those suffering from Kral's⁸ "benign senescent forgetfulness" (BSF) or Crook's⁹ "age-associated memory impairment" (AAMI), both of which advance very gradually, if at all, and early AD is not very clear. In a Cambridge field study¹⁰ about half of those with mild memory impairment later developed dementia, but it would have been hard to predict which. Among BSF subjects at the Maudsley Memory Clinic¹¹, however, there was a tendency for men and

those whose forgetfulness was noticed by others subsequently to develop dementia.

ORGANIC DISORDERS

The start of AD is usually manifest in memory impairment:

1. Things are mislaid at home, cannot be found in their once familiar places, or are left behind at home, in shops or in cars, buses and trains.
2. There is an increasing need to check that things have been done and reliance on aides-memoires; even so, appointments and plans are forgotten.
3. The same remarks are made and the same questions asked again and again, and conversation rambles on irrelevantly.
4. Recent information and activities are forgotten and messages are not passed on.
5. The start of a story is forgotten before it is over, so it is difficult to follow plays, films, books or news.
6. New locations, as when on holiday, are not learnt easily, and the patient may get lost.
7. Some things may be done twice over, like feeding an animal or cleaning teeth, and others not at all, like paying bills or taking needed medication for, say, heart-failure or diabetes.

At a later stage:

8. Once-familiar faces and places and locations seem less familiar, and eventually the sufferer may be lost in his/her own neighbourhood or even in his/her own dwelling.
9. The nearest and dearest may not be recognized, and treated as strangers, while true strangers may be greeted warmly as old friends or family.
10. The day, and the time of the day, are forgotten, and the patient may go shopping in the middle of the night, and be unable to find his/her own home on returning.
11. Sometimes the patient no longer knows his/her age or birthday, or that his/her parents are no longer alive, and has indeed entered upon a "second childhood".

Language impairment is usually regarded as a later feature of AD¹². The sequence of deterioration begins with tasks, such as naming, which use the semantic system, concerned with the meaning of words. There follow deviations and simplifications of syntax (grammar), and then phonemic breakdown (disordered use of sounds)¹³. One study¹⁴ found more deficits on the Boston Naming Test than the Mini-Mental State Examination in early

dementia, suggesting that naming difficulties may precede memory failure. Whether the common difficulty in finding proper names experienced by the over-50s¹⁵ is related to the later onset of dementia, or is merely a manifestation of "AAMI", has yet to be determined. Perseveration in AD is much less common than confabulation. Difficulties in understanding, as well as in finding, some words increase during the dementia, until finally there is almost total incomprehension and incoherence. Difficulties in reading and writing may precede those affecting the spoken word.

Loss of intellect is traditionally demonstrated by psychometric tests, which show a greater impairment of "performance" than "verbal" IQ¹⁶. It is also exposed by difficulty in defining concepts and explaining similarities in the course of cognitive testing. In everyday life it is mainly manifest in illogical thinking and often inconsistencies, e.g. "I live with my mother". "How old is she?" "Oh, in her 80s". "But how old are you?" "I'm 82". "Then you're about the same age as your mother?" "That's right!" There is a failure to draw appropriate deductions from environmental cues: a shopping expedition is undertaken in pitch dark, or, despite the blizzard evident outside the window the season is stated to be summer. A lack of "common sense" contributes to the increasing dependency.

Judgement and creativity are early casualties of dementia, and a good sense of humour may be the first loss. Taste in music, art, reading, clothing and decor may be coarsened, so that the patient wears a garish or incongruous garment, comes back from the shops with a tatty ornament which clashes with the fastidiously appointed living-room, only reads familiar books or gives up *The Times* for a tabloid newspaper. Exploitation by unscrupulous callers, who offer a few pounds for valuable heirlooms, is all too easy. Wills are changed without due regard for those who might have expected to inherit, in favour of some *parvenu* opportunist, causing deep hurt, disappointment and disputation when the patient has died. Judges and physicians become erratic and unreliable and artistic activity ceases or becomes facile, empty and repetitive.

Agnosia contributes, with amnesia, to disorientation. A failure of recognition of faces (prosopagnosia), places and objects bewilders the patient and his/her carers. Apraxia presents as difficulties in dressing—clothes are put on, if at all, in the wrong order, back to front and upside down—and feeding: knife and fork may have to be replaced by a spoon, and the patient may then use his/her hands or lap food from the plate. Apraxia may affect walking, when there is difficulty in judging the height of steps, or a change in the covering or the colour of the floor may be perceived as a step.

A personality change is not inevitable in AD, neither is it always for the worse. Occasionally, those who recognize their limitations and the need for others' help become less dominant and assertive and more docile, mellow and biddable. Commonly, however, as the dementia progresses the range of responses narrows, animation and spontaneity are subdued into apathy and indifference; the unpleasant label "vegetable-like" can apply. Some people become uncharacteristically coarse and disinhibited, swearing and using obscenities. There may be frequent, noisy, seemingly insatiable demands. Irritability, reproaches and angry outbursts can devastate and mystify carers. Regression in dementia facilitates the crude use of mental mechanisms, notably denial and projection: problems are denied or blamed on others. "There's nothing wrong with my memory—why, I can remember years back. It's other people who muddle me up by trying to catch me out. If they'd just leave me alone I'd be all right". A loss of insight, however, is not inevitable in AD, or not until loss of language prevents its expression. Alzheimer's original patient sometimes declared that she could not understand anything, and many sufferers are similarly painfully aware that there is something terribly wrong with their health.

Disordered behaviour is often the most distressing and challenging aspect of progressive AD¹⁷. Behavioural and psychological symptoms of dementia (BPSD) are strong predictors of caregiver burden and psychiatric morbidity¹⁸. Some forms arise understandably from personality change and cognitive deficits, while others might be core features of the dementia¹⁹. Studies based on those patients or clients referred to health and social services are likely to find disordered behaviour more often than in those who manage or are managed without such help at home, although few epidemiological studies assess behaviour as well as cognitive deficits. O'Connor *et al.*²⁰ found that among community residents in Cambridge, UK, such disturbed behaviour as demanding attention, repeating questions, using bad language, noisiness, temper outbursts, physical aggression and nocturnal wandering increased with the severity of dementia: 7% of those with mild disorder were aggressive and 42% of the severely demented.

Withdrawal and reclusiveness can have a protective function for those who find that the complexities of their former life are now beyond them. Self-neglect and squalor may be the inevitable consequence of incompetence and lack of help, often because it is refused, although sometimes demented people who live with others are reluctant to wash themselves or to change their clothes when it is time they did. Hoarding and clutter may result because of difficulty in taking the decision to throw rubbish away. Leaving the gas on, carelessness with cookers and inadvertent fire-raising are simple, although dangerous, consequences of forgetfulness and impaired judgement. Verbal abuse and acts of aggression are consistent (although still alarming) with explosive irritability, and the "catastrophic reaction" to exposure of cognitive incompetence in those who robustly deny it makes some sense, both as a form of protest and as a defence. Disinhibition—social, being excessively outspoken, tactless, fulsome or critical, yawning and nodding off during a conversation or a meal and departing from a gathering prematurely and abruptly; minor shoplifting (although this may be partly due to dysmnnesia); and, occasionally, sexual disinhibition, taking the form of stripping, exposure, open masturbation and importuning—cause carers and companions various degrees of embarrassment and shame. Noisiness may be associated with boredom, deafness, the desire for attention or depression (see below). Interfering with appliances and destructiveness may be a perverted form of "do-it-yourself"! Strange actions like talking to the reflection in the mirror as if it were another person and to photographs as if they were real people may be partly attributable to prosopagnosia.

Wandering and incontinence, two of the most troublesome behaviour disorders, are often overdetermined. Wandering may arise from utter boredom and underactivity; it is difficult to interest someone who can no longer read or enjoy television or radio and who is discouraged from going out alone for a walk or to the shops, lest they should get lost. Wanderers might have been in the habit of taking a walk with the dog or to buy cigarettes or a paper, at that time of day. They may be seeking their home even if actually at home; it may not be recognized as such, and there is a vain, vague, poorly (if at all) articulated quest for the home where they used to live. Wandering is sometimes a sign of agitated depression (see below) and occasionally a wanderer may be trying to get away from a loaded bowel! Restlessness towards nightfall ("sundowning") and nocturnal roaming²¹ by patients who have no idea of the time and are bored as well as restless, are a particular strain on carers²².

Incontinence of urine may occur for other reasons than dementia—stress incontinence, the frequency of a urinary tract infection, polyuria from diabetes or a prescribed diuretic. The foresight to urinate when the chance arises may have been lost, the location of the lavatory forgotten, or a vaguely similar place (like a cupboard or a storeroom) used in its stead. In advanced dementia

the patient may not appear to care any more where he/she excretes; alternatively, cortical control of the bladder may have been lost. Incontinence of faeces may be associated with impaction, compounded by poor hygiene, resulting in soiling and smearing²³.

The psychiatric features of the disease, described first by Alzheimer² and reaffirmed recently by Berrios and Brook^{24,25} and Burns and colleagues²⁶⁻²⁹, include affective disorder, paranoid and other delusional states and hallucinations. The affect in the disease is often blunted or labile, but it is now apparent that, at least among those seen by psychiatrists, depressive symptoms are common—63% of the 178 patients in Burns' series²⁸ having at least one. Depression was associated with less cognitive impairment, suggesting that it was related to insight. As most demented patients cannot describe how they feel, the Cornell Scale for Depression in Dementia³⁰ combines observed and informant-based signs of such possible depressive phenomena as diurnal variation, anorexia, insomnia, agitation and retardation and lack of engagement in activities, as well as overt misery.

Paranoid delusions of theft and intruders are a not uncommon reaction to losing things and are an example of projection. Such accusations are grievous to a loyal carer, as are those arising from morbid jealousy. Capgras' syndrome³¹, in which it is claimed that a stranger has adopted the appearance of someone well known to the patient, is rare but intriguing. Sixteen per cent of those in Burns' series²⁶, and 37% of those in Berrios and Brook's²⁵ were deluded.

Visual hallucinations tend to occur towards nightfall, and often take the form of little people—children, dwarfs or Lilliputians—who have to be fed. This is one reason why demented people may prepare meals for a household of people when only one or two are needed (another being because they have forgotten that their children have grown up and left home, as in Margaret Forster's moving novel *Have the Men Had Enough?*³²). Sometimes the very small people seem to emanate from a television set, which may act as a hallucinogen³³. Visual hallucinations are more common when the eyesight is impaired, and may be symptomatic of sub-acute delirium, superimposed on the dementia. Burns²⁷ found that 13% of his series of patients had visual hallucinations, and 10% auditory, but in clinical practice the latter are of little or no importance.

Finally, there are a number of neurophysical and other disorders associated with AD, including (towards the end of the illness) parkinsonism³⁴, gait apraxia³⁵, fits³⁶ and primitive reflexes^{37,38}. Weight loss in AD is particularly interesting, and not solely explicable by inadequate diet because of reluctance or forgetting to eat or to overactivity³⁹.

The course of AD is one of progressive deterioration from its insidious onset until death, 1–20 years later. Although younger patients live longer, they show the highest excess mortality⁴⁰. Measures of deterioration, like the Global Deterioration Scale⁴¹, rely upon a more uniform progression than is always found. Plateaux, when the clinical disorder seems for a while to be stationary, are not inconsistent with the diagnosis of AD⁴². Early language deficits⁴³ carry a poor prognosis for survival. Bondareff⁴⁴ postulated two forms of the disease: AD1, in older subjects, with a gradual course, mainly affecting memory; and AD2, in younger people, affecting temporoparietal functions and running a relatively rapid course. The two forms are also distinguished by their neuropathology and the pattern of neurobiochemical abnormalities. Death, when sooner or later it comes, is commonly from bronchopneumonia⁴⁵, developing in a person who by now is very helpless (often requiring to be fed) and debilitated.

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Assessment and Management of Behavioural and Psychological Symptoms of Dementia (BPSD)

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ASSESSMENT

The evaluation of BPSD should include: (a) a clear description of the target behaviour, including its antecedents and consequences (“ABC” behaviour analysis); (b) a search for external (e.g. physical environment, carer behaviour), physical (e.g. pain, delirium) and mental (e.g. psychosis) precipitating factors; and (c) an assessment of the risks posed by this behaviour to the patient, carer or other residents.

One good way to understand how to assess the range of BPSD is to use one of the many scales that have been developed^{1,2}. These tend to have a number of features in common. First, the principal carer should be the main source of information about the patient’s behaviour. Second, the frequency of the phenomenon in question may be a more useful question than “how severe is the problem?”, since the latter is more subjective. Third, the time frame should be specified—usually a month or a week.

The term “agitation” is widely used in research and clinical practice but has been used to denote restlessness or aggression or mood changes, each of which should be individually described when analysing the “agitated” patient. “Psychosis” should similarly be dissected into component phenomena: visual and auditory hallucinations, delusions and misidentifications of various types. The issue of which symptoms tend to co-occur is controversial but important, since such clusters may reflect distinct underlying aetiologies³.

MANAGEMENT

Until recently, there were few good quality trials of treatments for BPSD. This is in part because such trials are difficult to do. Patients are by definition most unlikely to be able to give consent to participate, “BPSD” are heterogeneous and staff are often reluctant to risk stopping medication, believing that they may be making patients easier to manage⁴. In trials of pharmacological interventions, “escape” or “rescue” medication (usually chloral or lorazepam) is needed. Nevertheless, the recognition of the burden these symptoms place on carers and of their tendency to result in the breakdown of caregiving in the home, coupled with the increasing evidence of harmful effects of conventional neuroleptics, has driven a fast-moving field.

Non-pharmacological Management

I consider the direct treatment by mere medicinal applications to be very limited. A kind and soothing reception, immediate removal of restraints, a warm bath, clean clothing, comfortable food, encouraging words, a medical treatment first directed to any manifest disease... A liberal diet, moderate use of malt liquor, exercise out of doors, employment, recreation, mental occupation, friendly intercourse and judicious religious attentions, are all important auxiliaries to amendment.

John Conolly, who was Medical Superintendent at Hanwell Asylum when he wrote this in 1847, was not writing specifically about dementia. However, there is no doubt that, overall, the management of BPSD requires more attention to psychosocial factors and to physical illness than to psychiatric factors. A variety of general supportive measures and specific behavioural interventions have been used.

General Measures

Dementia therapies. Reality orientation, reminiscence therapy and validation therapy are the best-known examples of therapies developed for use with elderly people. Although these interventions may provide valuable stimulation and foster therapeutic optimism, there is little evidence to suggest that they produce significant or sustained improvements in behaviour problems.

Stimulation-orientated therapies. Inadequate stimulation appears to have a negative impact on cognitively impaired elderly people. Training carers to identify and increase the number of pleasant activities for their dependants improves depressive symptoms⁵. Musical stimulation reduces disruptive vocalization in some patients⁶. For nursing-home residents, therapeutic activity programmes, as well as the physical environment, are important sources of stimulation. Rovner *et al.*⁷ found that a programme which included music, exercise and crafts, combined with guidelines on the use of psychotropic medication, reduced the prevalence of behavioural disorders and antipsychotic drug use. In one of the few environmental studies, Cohen-Mansfield and Werner⁸ found that simulating a home or natural outdoor environment within a nursing home using visual, auditory and

olfactory stimuli resulted in a modest reduction in pacing and other agitated behaviours.

Behavioural Interventions

Behavioural models, most commonly based on operant learning theory, have been used to design interventions for specific behavioural problems in dementia. An “ABC” analysis may suggest the implementation of specific behavioural techniques, including: (a) antecedent control—modification of events which trigger disruptive behaviour; (b) extinction—target behaviour is discouraged when it is not rewarded; (c) differential reinforcement of other behaviour (DRO)—the provision of positive reinforcement only for behaviours incompatible with the problem behaviour.

Personal care, for example, is a common antecedent to aggressive behaviour. This may occur because an intrusion into a patient’s personal space is misinterpreted as a threat. In such circumstances, careful prompting by the carer, an adequate explanation of the procedure and simple one-step instructions may reduce the perception of threat, and subsequently eliminate or reduce the defensive response. Similarly, if a nursing home resident’s only input from staff occurred following verbal outbursts, the staff response might serve to reinforce the behavioural problem. A behavioural analysis would allow staff to modify the antecedent (inactivity), or their own responses, through the use of extinction or DRO.

What is the evidence for the efficacy of behavioural interventions in dementia? Teri *et al.*⁹ have found that family carers can learn behavioural management strategies and that these interventions can reduce a wide variety of behavioural problems. In a controlled trial of a comprehensive carer training programme, which focused on behaviour management training and family support, carers’ negative reactions to behavioural problems were reduced, even though there was no reduction in the incidence of disruptive behaviours¹⁰. The majority of studies evaluating behavioural interventions in agitated nursing home residents have reported favourable results, mainly through the use of antecedent control and reinforcement of positive behaviours¹¹.

However, two larger recent studies failed to find a significant effect of behaviour management programmes^{12,13}.

Conclusion

Although psychosocial interventions have been shown to reduce behavioural disturbances, significant obstacles exist to their widespread implementation. There has been insufficient evaluation of clearly defined strategies targeted at specific behavioural problems. Therapeutic gains, in many cases, persist only for the duration of the intervention. In addition, staff compliance with behavioural strategies is often poor, perhaps unsurprising when their inadequate training and poor working conditions are taken into consideration. The successful implementation of psychosocial interventions in dementia requires the continuing support and training of professional and family caregivers.

Pharmacological Interventions

In this chapter, recent randomized studies are summarized and the effect of different drugs on clinically important measures is compared using “numbers needed to treat” [NNT = 1/(difference in proportion responding to each treatment)]. The lack of a consistently applied definition of “clinically important measures” can limit the validity of such comparisons, particularly when using data from meta-analyses¹⁴. Nevertheless, NNTs and NNHs (numbers needed to harm) are powerful and simple ways of

expressing clinically meaningful information, and are the best statistics available for making comparisons¹⁵. An NNT of less than 5 on a clinically important variable would usually indicate a drug that clinicians would find very useful.

“Typical” neuroleptics

Three meta-analyses have considered the efficacy of typical neuroleptics. In the first, Schneider¹⁶ showed an 18% difference between placebo and active drug in the proportion of patients responding to drug and placebo, equivalent to an NNT of 5.6. A more detailed presentation of 16 randomized trials since 1966 found an overall NNT for “improvement” on neuroleptics of 3.8 and an NNH of 4.0¹⁷.

Thioridazine. The Cochrane meta-analysis of the effect of thioridazine in dementia¹⁸ found only one large placebo-controlled study that reached inclusion criteria¹⁹. This was a multicentre study of 610 institutionalized patients in which 4 weeks of thioridazine was compared to placebo and diazepam. Thioridazine improved the “anxiety/tension/fears/insomnia” factor of the Hamilton Anxiety Scale with an NNT of 2.6, and the “cognitive impairment/agitation/depressed mood/behavioural change” factor with an NNT of 3.1. However, the overall conclusion of the Cochrane authors was that, because of the poor quality of reporting of trials: “If thioridazine were not currently in widespread clinical use, there would be inadequate evidence to support its introduction . . . [It] has minimal or no effect on global ratings, while other drugs such as chlormethiazole are superior to it on behavioural ratings”.

Haloperidol. Devanand and colleagues²⁰ (1999) addressed the question of optimal dosing of haloperidol. This was a study of 71 outpatients with Alzheimer’s disease with a mean Mini Mental State Examination (MMSE) of 10. The majority of patients showed both psychosis and disruptive behaviours, so it was not possible to tease out any differential effect on the two classes of disorder. Response rates according to three sets of criteria were greater with the standard dose of 2–3 mg daily (NNT 3.3–4.0) than with low dose (0.5–0.75 mg). A subgroup (20%) of those on 2–3 mg daily developed moderate to severe extrapyramidal signs (NNH = 5), but did not meet consensus criteria for dementia with Lewy bodies. The authors recommend a starting dose of 1 mg/day with upward titration.

Atypical Neuroleptics

Risperidone. The largest good quality trial to date of drug treatment for BPSD has been that of Katz *et al.*²¹. Nursing home residents with dementia were treated for 12 weeks with either 0.5 mg, 1 mg or 2 mg risperidone daily, or placebo (approximately 150 in each arm of the study). Most had severe dementia: the mean MMSE was 6.6. The primary outcome measure was a 50% reduction in the BEHAVE-AD, which occurred in 33% taking placebo, 45% on 1 mg (NNT = 8.3) and 50% on 2 mg (NNT = 5.9). Interestingly, risperidone appeared to have an anti-aggression effect (NNT = 3.4–3.9) that was independent of its effect on psychosis (NNT for delusions 7.1 at 1 mg, 10.0 at 2 mg) or its tendency to cause somnolence. The dropout rate varied from 27% in the placebo group to 42% in the 2 mg group. The NNH for sedation was 11.2 for the 1 mg dose and 5.0 for the 2 mg dose. Extrapyramidalism was more common at 2 mg (NNH = 7.2) than at 1 mg (NNH = 18.5). The optimal dose of risperidone is therefore 1 mg daily.

Olanzapine. Street *et al.*²² studied 206 nursing home patients with a mean MMSE of 7.3 in a trial of six weeks of olanzapine or placebo. The main outcome measure was a 50% reduction in the Neuropsychiatric Inventory (NPI) score for a newly defined cluster of symptoms (hallucinations, delusions, agitation). The optimal dose (5 mg) was as effective as typical neuroleptic (NNT=3.3) but seemed to be associated with relatively high rates of adverse reactions. The response rates (66% on 5 mg, 52% on 10 mg, 43% on 15 mg and 36% on placebo) suggest that doses higher than 5 mg are likely to be suboptimal. Somnolence and gait abnormality were the commonest side effects (NNH = 5.4 and 5.7, respectively) This study illustrates the difficulties that occur in comparing adverse drug reactions (ADRs) across studies, since it is very likely that ADRs were more assiduously sought in this study than earlier studies of typical neuroleptics. The lower dose of 2.5 mg, which is now available—and which one would expect to be associated with fewer ADRs than 5 mg—was not tested.

Carbamazepine. Few trials have published significant differences in responder rates on measures of Global Clinical Impression (CGI). One exception is the study of Tariot *et al.*²³. At least minimal improvement on the CGI was apparent in 77% of those taking carbamazepine compared to 22% on placebo (NNT=1.8). Furthermore, the reduction in the number of cases requiring “a great deal” or “almost constant” extra nursing time to deal with the agitation was 37% in those taking carbamazepine and 8% in those taking placebo (NNT=3.4). This was a relatively small (ca. 25 subjects in each arm) 6 week study of predominantly female nursing home residents with severe dementia (mean MMSE 6.0), 92% of whom were aggressive. Carbamazepine was started at 100 mg/day. The modal dose reached was 300 mg/day. The NNH for any ADR was 3.3, although these were considered clinically significant in only 2 of 16 cases. The large effect size may have been at least partly attributable to the fact that a non-blind physician adjusted the dose of carbamazepine—a strategy which, of course, parallels the clinical situation, but is not commonly employed in more recent studies.

Trazodone. Trazodone is widely used for BPSD, although there has only been a single small randomized trial of its effectiveness, in which it was shown to be comparable in effect to haloperidol²⁴. The CGI was “much improved” or “very much improved” at week 9 in 57% of cases on haloperidol and 71% on trazodone. However, trazodone was associated with fewer side effects (14%, mean dose 218 mg) than haloperidol (50%, mean dose 2.5 mg). Interestingly, post hoc analyses suggested that those who were verbally aggressive or had repeated behaviours or mannerisms were more likely to respond to trazodone. Those who paced, were generally restless or made unwarranted accusations were more likely to respond to haloperidol. This raises the possibility of two syndromes of behavioural change in dementia, one associated with abnormalities of dopaminergic function and one associated with serotonergic dysfunction. The replicated finding that those with depression early in dementia are more likely to go on to be physically aggressive is also consistent with the possibility that there is a distinct serotonergic syndrome of behavioural change in dementia^{25,26}. It is not necessary for other signs of depression to be present to justify a trial of an antidepressant for aggressive behaviour.

Xanomeline. Xanomeline is a cholinomimetic with agonist activity at postsynaptic muscarinic M1 and M4 receptors. Bodick *et al.*²⁷ presented data from a large (87 in each arm), 6 month study of patients with mild to moderate AD. Three doses of xanomeline were compared with placebo. The dropout rate was high, ranging from 35% in the placebo group to 59% at the highest dose of active drug (NNH = 4.2), precluding its further use in the formulation used. However, there was a dose-related

response to xanomeline of hallucinations (NNT=2.1 at the highest dose) and a variety of other behaviours (e.g. vocal outbursts, delusions and mood swings; NNT=2.7–6.4).

Rivastigmine. The cholinergic nucleus basalis of Meynert is in double jeopardy in patients with dementia with Lewy bodies (DLB) because it is likely to degenerate as a result of both Lewy body and neurofibrillary change. Cortical cholinergic parameters show even more marked depletion in DLB than in AD. There is now evidence showing that patients with DLB are preferential responders to cholinesterase inhibitors. McKeith *et al.*²⁸ conducted a 20 week placebo-controlled study of 3–12 mg rivastigmine in 120 patients with probable or possible DLB of mild to moderate severity. The main outcome measure was a 30% reduction on a newly defined cluster of symptoms derived from the NPI: hallucinations, delusions, depression and apathy. A 30% reduction was chosen because it was comparable to the effect size seen with neuroleptics. The main result was based on the observed case analysis: 63% of those on rivastigmine responded, compared to 30% on placebo (NNT=3.0). Although there was no overall difference in the number of ADRs (59 vs. 61%), those taking rivastigmine were significantly more likely to drop out (31% vs. 16%; NNH=7.1). As well as improvements in the pre-defined cluster, there were also marked improvements in a computer test of attention in those on rivastigmine.

These two studies provide good evidence that hallucinations in particular, and possibly other BPSD, respond well to cholinergic agonism.

When to Consider a Trial without Medication

Neuroleptics have unpleasant side effects. Sedation, falls, hip fractures and tardive dyskinesia are common sequelae. Indeed, the prescription of typical, rather than atypical, neuroleptics is likely to be a false economy, particularly if there is any risk that the drug will be taken for more than a brief period. Trials without neuroleptics are a legal requirement in nursing homes in the USA, following the OBRA-1987 regulations. What is the optimal duration of treatment before a trial without medication is indicated?

Since evanescent symptoms do not require long periods of treatment, an important related question is “How often would the symptom normally be expected to resolve without drug treatment within a given time?” Levy *et al.*²⁹ assessed 181 outpatients and found that 60–70% of those with agitation or psychosis still had the symptom 3 months later. Persistence was more common amongst those with low MMSE scores (12–17). Devanand *et al.*³⁰ in an outpatient study of 235 patients with early AD, found that 74% of those with wandering or agitation had the symptom 6 months later. Comparable figures for other behaviours were: physical aggression 53%, hallucinations 52%, paranoid ideas 45%, misidentification 56%. In a group of 48 patients with autopsy-confirmed AD, Hope *et al.*³¹ found that aggression and hyperactivity were both still present in 75% of cases after 8 months, and hallucinations were still present in 75% of cases at 11 months. All these studies were potentially directly or indirectly confounded by the effect of psychotropics on the natural history of behaviour. Nevertheless, these data suggest that BPSD may endure for up to 8 months in approximately 75% of cases and be evanescent in 25%.

How often will trials without treatment succeed and how often will they fail? Cohen-Mansfield *et al.*⁴ studied the effect of withdrawal of drugs from nursing home residents, many of whom will have had dementia. The doses of haloperidol (mean=0.9–1.3 mg), thioridazine (25–27 mg) and lorazepam (0.7–0.9 mg) that were being used were within current standards of care. The design

was a cross-over study with 6 weeks in the drug or placebo arm. Of 194 residents who were taking regular medication at the start of the study, only 58 (30%) participated, but only 16 (8%) cases were excluded for reasons that potentially related to their behaviour. Thirty-five (60%) completed the study. Dropouts due to emergence of BPSD were not more significantly likely to occur in the withdrawal phase (six cases) than the continuation phase (three cases). After correction for multiple comparisons, BPSD were not more likely to re-emerge following drug withdrawal than with continuation. There was also a slight improvement in cognitive function. At face value, this study suggests that a trial without medication at 6 weeks is indicated. However, the difficulty of sustaining non-pharmacological approaches to the management of BPSD in nursing homes was shown by the fact that, during the 22 weeks following discontinuation of the original drug, less than one-third of cases remained free of psychotropic medication. The mean time until receiving another psychotropic drug was 21 days.

SUMMARY

Old age psychiatrists are often called for advice when psychosocial interventions have been tried and have failed, or the resources of the staff to try such interventions are limited. Careful judgement is needed in these circumstances, since a decision not to prescribe can make a precarious situation worse. Drug treatment can provide a breathing space during which "something" has been seen to be done, as well as having a useful "real" impact in the 75% of cases in which the behaviour persists. The relatively high placebo effect (often 30–40%) can be useful, as long as the maxim "primum non nocere" is borne in mind. It is important not to leave patients on medication without having a trial of drug withdrawal. Recent demonstrations of the efficacy of atypicals, carbamazepine, antidepressants and cholinomimetics are important evidence that this most neglected area of psychiatry need not be dominated by therapeutic nihilism. Brief programmes of behavioural management are probably only effective briefly, if at all. Continuing support and training are required for successful psychosocial interventions.

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Eating Disorders in Alzheimer's Disease

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Eating disorders are a recognized feature of dementia, which include a preference for sweet foods (11%), increased (21%) or decreased (22%) consumption and eating non-food substances (3%) and are particularly common in Alzheimer's disease (AD)¹. Clinical¹ and subclinical³ swallowing problems, general dental health and oropharyngo-oesophageal function are all important. The basic necessity of eating to maintain health, the frustration that disordered eating causes to caregivers and the increased risk of institutionalization and death all make this an important topic⁴.

A number of rating scales to evaluate psychopathology in dementia now include an eating disorder subcategory (e.g. Neuropsychiatric Inventory), whilst other scales have been developed specifically to assess eating disorders in AD⁵.

Several neurotransmitters have been linked to eating disorders in AD. Reduced neuropeptide Y and norepinephrine are associated with anorexia, whilst the action of galanin in the hypothalamus is thought to increase fat intake and impact upon cholinergic hippocampal systems.

Studies have shown associations between specific brain changes and eating disorders in AD, e.g. hyperorality with widening of the third ventricle and frontal and occipital lobe atrophy, Klüver-Bucy syndrome features with temporal lobe atrophy¹⁴ and low body weight with temporal cortical atrophy⁶. Hyperphagia may be associated with increased calorific need in patients with motor restlessness, whilst younger people with more severe dementia who are not restless may over-eat because they respond to any food stimulus, possibly as a manifestation of frontal lobe pathology⁷. The European Commission has focused upon weight loss in AD⁴.

In practice there is little evidence to inform the management of these problems. The best approaches generally involve common sense and clinical judgement. The most frequent clinically significant problems relate to poor appetite and weight loss. A first step is to assess and treat underlying disorders. This may vary from assessment of oral health to the pharmacological treatment of a concurrent depression. Educating care staff and informal carers about some of the changes in food preference and encouraging flexibility with the content and timing of an individual's diet is often an effective remedy⁸⁻¹⁰.

Giving a diet with a higher proportion of sweet foods, such as desserts, chocolate, cakes and biscuits, is a pragmatic approach. This is helpful as it takes pressure off carers to produce a strictly balanced diet of cooked dinners, which eases the stress of mealtimes. Other types of difficulty are less frequent, and are

probably best treated after a detailed individual evaluation, using techniques such as an Antecedent-Behaviour-Consequence ("ABC") diary. Work pertaining to ideational apraxia¹¹ and attribution theory¹² has contributed to the management of these problems. The management of eating disorders in AD raises important ethical issues¹³.

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Pathology of Vascular Dementia

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Cerebrovascular disease is usually considered to be the second commonest cause of dementia, although the precise prevalence and the frequency of combination with Alzheimer's disease (AD) is very variable. In the West, approximately 15% of autopsy-studied cases of dementia have been attributed to vascular disease alone and 8–18% to a combination of vascular disease and AD^{1,2}. However, this has been challenged in a recent autopsy study of patients in a dementia clinic, where dementia could not be attributed in any case to vascular disease alone³. Vascular dementia is commoner in the Far East than in the West and still remains the commonest form of dementia in Japan⁴. In the West, it may also be commoner than AD in the very elderly⁵. Estimates based on death certification and clinical diagnosis without pathological confirmation are liable to be significantly inaccurate^{6,7}.

If it is difficult to determine how frequent vascular dementia is, it is also difficult to define what it is. For example, the terms “arteriosclerotic” and “multi-infarct” dementia are often used synonymously and these, together with a lacunar state and Binswanger's disease, have been grouped together⁸. In practice, three separate forms of vascular dementia can be identified, although there is often overlap between these: multi-infarct dementia; dementia due to single strategically-placed infarcts; and dementia due to diffuse white matter damage⁹.

MULTI-INFARCT DEMENTIA

Multiple areas of infarction involving both cortical and sub-cortical locations, separated in both time and space, may be associated with dementia, although in its pure form this is probably uncommon¹⁰. The location of the infarcts is probably more important than the volume of tissue lost (100 ml is the oft-quoted figure)^{1,11}. Dementia may occur with relatively little tissue loss, particularly if infarcts are strategically located (see below) and the effect of subsequent infarcts may be synergistic rather than additive¹².

Multiple small infarcts of deep white matter or grey structures (“lacunar state”) may also be associated with dementia, although this often overlaps with dementia due to diffuse white matter damage¹³. Patients with a pure lacunar syndrome show features of a subcortical dementia—psychomotor slowing, poor concentration, indecision and apathy—without features of cortical dysfunction⁹. Hypertension is the most important risk factor for dementia of this kind and that is particularly true for dementia due to multiple lacunar infarcts^{13,14}.

DEMENTIA DUE TO SINGLE STRATEGICALLY PLACED INFARCTS

Dementia may result from single infarcts involving certain areas of the brain, such as the angular gyrus of the dominant parietal lobe, the medial thalamic nuclei and head of the caudate nucleus, especially if bilateral, the globus pallidus, basal forebrain and hippocampus^{9,15}. The cognitive effects of infarction of the angular gyrus may closely resemble AD, although the sudden onset of the cognitive deficit may point to the correct diagnosis¹⁶.

DEMENTIA DUE TO DIFFUSE WHITE MATTER DAMAGE

Whilst white matter damage due to multiple infarcts may be the substrate of dementia, there are also forms of more diffuse white matter damage due to small-vessel disease which result in dementia, and these are probably commoner than dementia due to single or multiple infarcts¹⁷.

Binswanger's disease is considered to be a well-established clinicopathological entity: ischaemic periventricular leuko-encephalopathy manifested clinically by subcortical frontal executive dysfunction, parkinsonism, urinary incontinence, mood changes and pseudobulbar palsy¹⁸. However, it is still the subject of considerable controversy regarding both its frequency¹⁹ and its relationship with the radiological entity of “leuko-araiosis”—the appearance of low attenuation areas in the white matter on CT scanning²⁰. CT abnormalities are common in the elderly and may not be associated with dementia^{21,22} and the clinical significance of leukoaraiosis remains incompletely defined²³. It certainly does not equate to Binswanger's disease, despite suggestions to the contrary²⁴.

A review of the pathological features of all the reported cases of Binswanger's disease in the world literature, 1912–1986, together with Fisher's review, reveals that the vast majority of cases designated as Binswanger's disease show consistent pathological changes^{25,26}. The general autopsy shows evidence of prolonged systemic hypertension. The brain is of average weight but there is diffuse dilatation of the lateral and third ventricles, with rarefaction, discoloration and a rubbery texture of the periventricular white matter, particularly in the occipital regions. The arteries of the Circle of Willis are affected by atheroma, which in 60% of cases is severe. Lacunar infarcts are usually present but one-third of brains also contain large infarcts. Microscopically, there is incomplete diffuse demyelination, with gliosis and microinfarcts affecting the periventricular white matter but sparing the subcortical fibres, corpus callosum, anterior

commissure and internal capsules. The walls of white matter penetrating arteries are hyalinized but the vessels are patent. This condition is probably caused by hypoxic-ischaemic damage to the distal watershed periventricular white matter, secondary to narrowing of arterioles^{23,27,28}.

A closely related cause of dementia due to diffuse white matter damage is CADASIL (cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy)²⁹. This is a disease that usually presents in the fifth decade with strokes or dementia, and is an inherited vascular disorder due to mutation affecting the *Notch3* gene on chromosome 19. Pathologically, the disease is very similar to Binswanger's disease, with widespread lacunar infarcts and diffuse cerebral white matter degeneration. The distinguishing feature is the presence of non-hypertensive arteriolosclerosis, characterized by deposition of granular osmiophilic material in relation to vascular smooth muscle cells, with degeneration of these cells and thickening of the vessel walls. This is a systemic vascular disorder, although other tissues are less severely affected than the brain, and the diagnosis is therefore possible on skin or muscle biopsy, although genetic analysis is required for confirmation³⁰.

In the experience of the author, vascular dementia is much less common than AD. However, it is given a significance disproportionate to its incidence by the fact that some causes of vascular dementia—such as giant-cell arteritis³¹ and thromboangiitis obliterans³²—are potentially treatable, and that vascular dementia as a whole is theoretically preventable by addressing risk factors for cerebrovascular disease³³.

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International Criteria for Vascular Dementia and Their Problems: ICD-10, DSM-IV, ADDTC and NINDS-AIREN

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There are two sets of criteria currently available for the diagnosis of vascular dementia. The *Diagnostic and Statistical Manual of Mental Disorders*, 4th edn (DSM-IV)¹ and the *Classification of Mental and Behavioural Disorders*, 10th Revision (ICD-10)² are general diagnostic tools and outline the criteria but do not operationalize them. The second set, the State of California Alzheimer's Disease Diagnostic and Treatment Centers³ and the National Institute of Neurological Disorders and Stroke and the Association Internationale pour la Recherche et L'Enseignement en Neurosciences (NINDS-AIREN)⁴ criteria, are developments of the first two and offer operational criteria.

There are three fundamental flaws and several lesser errors lying in the details of these criteria. All the criteria are fundamentally similar in that they first require the presence of dementia, which is based on Alzheimer's disease (AD)-type features, and subsequently identify vascular dementia as a subset of all dementia, using vascular features that are often operationalized using the ischaemic score⁵.

Unfortunately, the use of Alzheimer-based criteria for the diagnosis of dementia has now been clearly shown to be wrong. Vascular dementia does not closely resemble AD, even when Alzheimer-based criteria have been used for case identification⁶. Thus, cases selected using current criteria represent only a subset of all vascular dementia. This is the first major flaw in the current criteria.

The second problem is the increasing recognition of mixed dementia. Originally, vascular dementia and AD were separated by the presence of large infarct volumes⁷. In the nun study, in which 47% of demented patients had mixed disease, very small amounts of cerebrovascular disease profoundly altered the age of presentation and speed of progression of what otherwise appeared to be AD⁸. Reclassification of autopsy data to allow small infarct volumes to convert a diagnosis of AD to mixed dementia increased the proportion of cases with mixed dementia from 2% to 18%⁹. Thus, mixed dementia is far more important than was realized when the current criteria, which scarcely recognize mixed disease, were prepared. No good method yet exists for separating mixed dementia. Criteria that first select cases that appear like AD and then subselect those with vascular features might form an excellent basis for doing so; unfortunately, this is what the current criteria for vascular dementia do and it is very likely that much of the reported data about vascular dementia is in fact about mixed disease.

The next fault is the level of severity. The criteria define dementia as the level of cognitive impairment at which normal daily functions are impaired, and therefore will identify only late cases, underestimating the prevalence of cognitive impairment due to vascular disease. More importantly, they prevent identification of early cases that would benefit most from preventative treatment¹⁴. This important early stage has been termed "vascular

cognitive impairment" (VCI)¹⁰. VCI is important because vascular disease is the largest single identifiable risk factor for dementia apart from age, and the only one currently treatable.

The above features constitute the fundamental faults of the current criteria. There are a number of additional problems that lie in their details. Differences in the details lead to great differences in the proportion of cases identified as having vascular dementia³.

Correction of these faults requires wholesale revision of the criteria with the development of new criteria based on data rather than supposition¹¹⁻¹³.

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Vascular Dementia

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EPIDEMIOLOGY

For many years, dementia was thought to be due to vascular disease. This seemed logical, largely on the basis that atherosclerosis and dementia are common in the elderly. In the 1970s and 1980s, many papers appeared stressing that vascular dementia was over-diagnosed¹, and most came to the conclusion that Alzheimer's disease (AD) was the commonest cause of dementia, even in patients who had had a stroke. In recent years the pendulum has swung a little back towards vascular disease being a more significant cause. Furthermore, the clear margins between some types of vascular dementia and AD have become blurred. Indeed, brain infarction may play an important part in determining the expression of the clinical symptoms of AD².

The epidemiology of vascular dementia is fraught with difficulties because of the lack of a reliable definition to act as a gold standard. Many groups have come to a different consensus about the criteria for the diagnosis (e.g. DSM-IV, ADDTC, NINDS-AIREN, ICD-10 and Hachinski score) Not only is the clinical diagnosis ambiguous, partly because many patients have mixed dementia (i.e. both AD and vascular disease), but the pathological differentiation of the varying types of dementia is also blurred. Does one then use clinical series, preferably community-based, or autopsy series? Reliable data are very sparse. Referral bias remains a major problem in hospital based series. In a community-based study in those aged over 85 from the USA, dementia was found in 30%³. In Europe and the USA, vascular disease accounts for 10–40% of all cases of dementia, whereas in Japan vascular disease is more common than AD; the lifetime risk of developing vascular dementia is approximately 25%^{4–6}. Pathological studies suggest that cases of mixed AD and vascular dementia may be as common as those with just vascular dementia.

Equally good reasons can be made for thinking that vascular dementia remains both underdiagnosed and overdiagnosed^{7,8}. Dementia is common following stroke occurring in approximately 16% after the first stroke⁹.

RISK FACTORS

The risk factors for vascular dementia have rarely been studied in isolation to stroke. The major risk factors are age, race, hypertension, diabetes mellitus, hyperlipidaemia, smoking, transient ischaemic attack and heart disease, especially ischaemic heart disease and atrial fibrillation¹⁰. The level of education, volume of cerebral loss, degree of cerebral atrophy and presence of periventricular white matter lesions are important in the devel-

opment of dementia¹¹. Other possible risk factors include polycythaemia, homocysteinuria, hyperfibrinogenaemia, alcohol excess and lack of physical exercise.

Most of these risk factors predispose to the atherosclerotic process in all its many guises (e.g. large vessel atherosclerosis, small vessel lacunae and white matter leukoaraiosis). However, there are a number of other causes for vascular dementia which are worthy of separate mention. In the last 10 years our understanding that genetic factors can predispose to vascular disease have gained increasing importance, with the detailed understanding of the families with cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL), mitochondrial diseases and hereditary cerebral haemorrhage with amyloid^{12–14}. CADASIL has been shown to be linked to chromosome 19: individuals are predisposed to migraine, vascular events and dementia¹². The genetics of mitochondrial encephalomyopathy with lactic acid and stroke-like episodes (MELAS) is of interest; only men are affected, with the disease being transmitted only by the maternal line. The clinical features include stroke-like episodes, lactic acidosis, short stature and deafness¹³. The diagnosis is confirmed by DNA analysis to look for point mutations in the RNA gene. Other rare causes of multi-infarct dementia and stroke include the anti-phospholipid syndrome^{15,16}, arteritis¹⁷ and late effects of cranial radiotherapy, amongst others^{18–20}.

CLINICAL TYPES

The pathological findings in vascular dementia are diverse. Findings range from large cortical infarctions (Figure 48.1) to small discrete deep infarcts (Figure 48.2) and to diffuse areas of white matter low-attenuation changes as seen on CT scan (Figure 48.3). Several of these changes may co-exist in the same patient (Figure 48.4).

Cortical Infarcts

Multiple cortical thrombo-embolic infarcts are a common cause of multi-infarct dementia. This will follow embolic occlusions from either the heart or the great vessels, primarily in the neck, e.g. carotid stenosis, or primary thrombosis in the major cortical arteries of the cerebral hemispheres, e.g. antiphospholipid syndrome^{15,16} and arteritis¹⁷. Dementia may rarely result from multiple watershed infarcts in the territories between the middle and posterior cerebral arteries in the cortex: hypotension is the usual cause secondary to prolonged cardiac arrest.

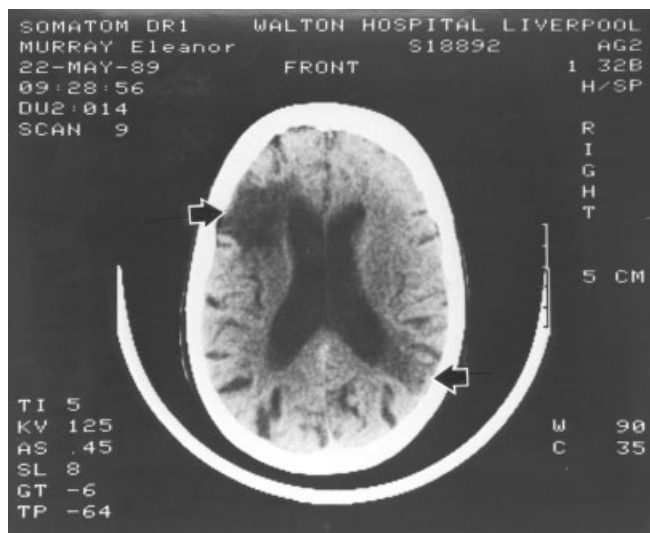


Figure 48.1 Multiple cortical infarcts

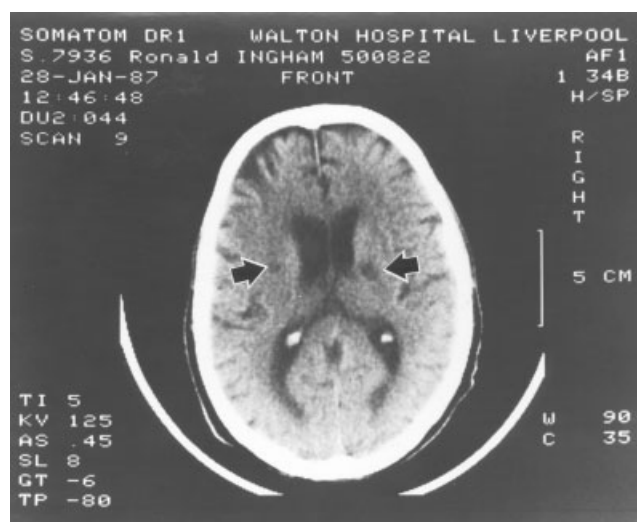


Figure 48.2 Multiple lacunae

A strategic single cortical infarct in the dominant angular gyrus can also cause a picture similar to AD with dysphasia, visuospatial disorientation, agoraphobia and memory loss. Multiple cerebral haemorrhage, which particularly occurs in cerebral amyloid, may cause dementia.

Lacunar Infarction

Lacunae are small deep sub-cortical infarcts or haemorrhages. A single lacunar stroke commonly presents with pure motor hemiplegia, pure hemisensory loss or ataxic hemiparesis²¹. Multiple lacunae (état lacunaire) was described by Marie to produce dementia, dysarthria, small-stepping gait, incontinence and emotional lability. Fisher doubted the existence of a multi-lacunar dementia, although there is now no doubt that vascular dementia does occur after multiple lacunar infarcts^{22,23}. Lacunar infarcts are common in hypertension and rarely follow embolic occlusion from the heart or major arteries.

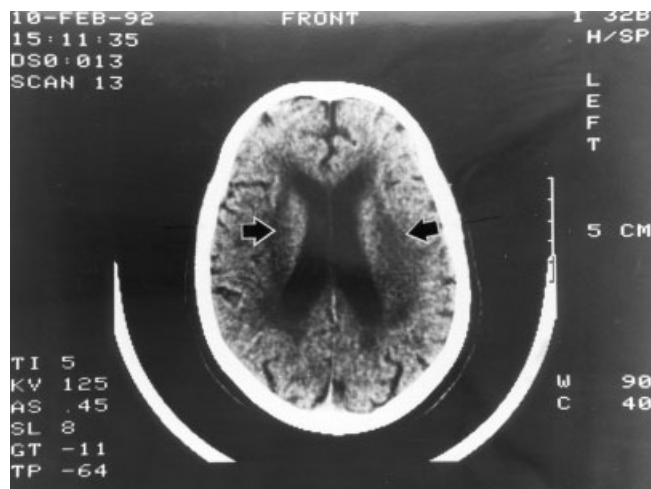


Figure 48.3 Leukoaraiosis

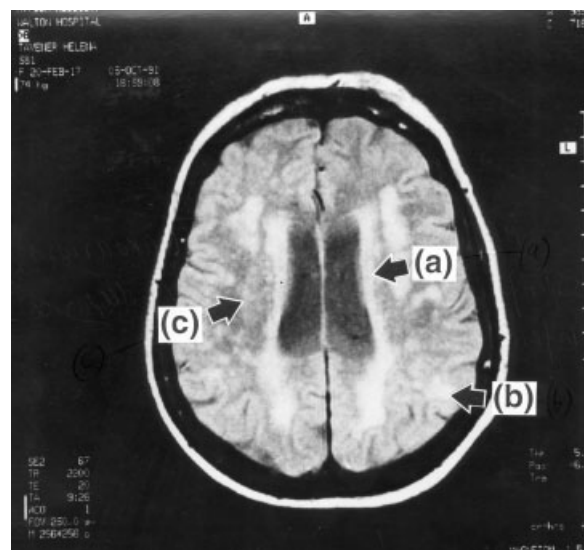


Figure 48.4 Cortical infarcts, lacunae and leukoaraiosis coexisting in the same patient

While multi-infarct dementia is nearly always secondary to multiple lesions, it is possible to see memory loss after a single infarct. Discrete thalamic infarction especially in the paramedian region, may present as memory loss, often associated with somnolence and eye movement disorders^{24,25}.

Leukoaraiosis

This is largely a radiological diagnosis describing the diffuse or patchy low-attenuation changes seen on CT or MRI scans in the deep white matter²⁶. Binswanger (1894) described eight patients with slowly progressive dementia who, at post mortem, were found to have softening and loss of deep white matter with compensatory ventricular enlargement. There is also gliosis, thickening of the arteries, within the abnormal white matter; many also show discrete lacunar infarction, although these are often too small to be seen on CT and MRI scanning. The

diagnosis of Binswanger's disease (subcortical arteriosclerotic encephalopathy; SAE) has become increasingly popular.

It was initially thought that these CT findings were specific for Binswanger's disease. However, similar CT findings can occasionally be seen in normal people or even those with AD. Because of this, the term "leukoaraiosis" (Greek, for white matter of loose texture) was used to describe the CT appearances²⁷.

Clinically there is a stepwise or progressive course, characterized by pseudo-bulbar palsy, cognitive, behavioural and gait disturbances, with focal neurological deficits in approximately 30%. The dementia tends to be subcortical in type. Many patients are hypertensive and the pathological changes are thought to be secondary to chronic ischaemia in the deep end arteries.

Although many patients are hypertensive, the final insult is probably due to hypoperfusion. Indeed, Sulkara and Erkinjuntti have reported acute dementia after periods of hypotension and cardiac arrhythmia; CT showed leukoaraiosis²⁸. Although hypertension needs to be treated, episodes of hypotension need to be carefully avoided, as vasodilatory reserve is impaired²⁹.

Inzitari *et al.* noted leukoaraiosis to be present in 100% of patients with multi-infarct dementia (MID) and 33% with AD compared with 11% of the control population, although even in this control population there was evidence of intellectual fall-off, but not severe enough to be labelled dementia³⁰. A history of stroke was four times more likely in those with leukoaraiosis than in those without; most, however, die from cardiac causes³¹. Overall there appears to be considerable overlap between leukoaraiosis, subcortical atherosclerotic encephalopathy and multiple lacunar states.

In CADASIL, the MRI usually shows a combination of leukoaraiosis and multiple lacunar infarction.

Cerebral Amyloid Angiopathy

Although cerebral amyloid angiopathy has been known about for many years, its role in the aetiology of dementia has attracted much attention³². Amyloid deposition in the small and medium-sized arteries and arterioles may present with spontaneous haemorrhage or thrombosis. More interestingly, there have now been many reports of cerebral amyloid angiopathy (CAA) presenting as dementia without any preceding history of stroke-like episodes. CAA is associated with areas of small subcortical infarction and histological changes similar to Binswanger's SAE.

Senile plaques in AD contain an amyloid core which appears to be identical to CAA. Amyloid plaque cores are often found immediately adjacent to amyloid-laden capillaries. CAA is present in more than 80% of cases of AD³³.

The association of AD with leukoaraiosis and cerebral amyloid angiopathy has renewed speculation that vascular risk factors may play a role in the pathogenesis of AD³⁴.

MANAGEMENT

Clinical Assessment

The following criteria point towards a diagnosis of vascular dementia:

1. Symptoms and signs of stepwise stroke-like episodes.
2. Vascular risk factors.
3. Evidence of vascular disease on CT and MRI.

The validity of the different scoring systems is discussed elsewhere (q.v.). Factors suggesting that the dementia is not vascular include:

1. Absence of the above.
2. Early presence of extrapyramidal and autonomic features.
3. Early hallucinations.
4. Cerebellar signs.

It should be noted that it is sometimes difficult in those who present with gait problems to distinguish leukoaraiosis/multiple lacunar strokes from normal pressure hydrocephalus and progressive supranuclear palsy: it is important, then, to look for poor upgaze, axial rigidity and perform CT or MRI scanning.

There is no single illness making up vascular dementia. The clinical management of any patient with vascular dementia should include a summary of the clinical presentation, the probable pathogenesis, risk factors and site of damage. This will allow a rational approach to investigation and medical treatment to minimize the risk of further recurrence/progression. An attempt should be made to decide whether the dementia is primarily a thrombotic or an embolic process from the heart or great vessels, a hypertensive deep white matter disease, a hypoperfusion process or even one of the rarer causes of stroke.

Investigation

Investigation should include a routine vascular screen and then further investigations, depending on the proposed pathophysiology. It is important to assess the routine vascular risk factors, especially as Meyer *et al.* have suggested that improved cognition may occur after the control of risk factors in MID³⁵. A careful assessment of the cardiovascular system is necessary, particularly paying attention to blood pressure and possible sites of emboli from the cerebral arteries (i.e. carotid stenosis) and heart, (especially valvular heart disease, atrial fibrillation or tumour and left ventricular thrombus) in those with cortical infarctions. The CT/MRI scan appearances in vascular dementia have already been discussed in some detail.

All patients should have a full blood count (for polycythaemia or thrombocythaemia), ESR (for arteritis), blood sugar, chest X-ray and ECG. I also check fasting lipids, as cholesterol is an important risk factor for ischaemic heart disease and most vascular patients die a cardiac death. Neurosyphilis remains treatable. Evidence of arteritis and the antiphospholipid syndrome should be looked for, both clinically and by checking various blood tests, such as CRP, complement levels, antinuclear factor, DNA binding, lupus anticoagulant and anticardiolipin antibodies. If the ESR is raised or the patient is over 50 it is important to consider the arteritides, such as granulomatous angiitis and temporal arteritis^{17,36}.

The diagnosis of granulomatous angiitis can be very difficult and may require both angiography and brain/meningeal biopsy. It is often difficult to know how often one should pursue these investigations, but a stuttering onset of recent vascular dementia with a raised ESR and abnormal lymphocytic cerebrospinal fluid (CSF) should alert the physician. Unfortunately, in some cases the ESR and CSF may be normal. A high index of suspicion is needed because with steroids and cyclophosphamide these patients can do very well^{17,20}. In spite of this, a raised ESR in stroke frequently remains unexplained.

Ultrasound scanning of the carotid arteries to detect carotid stenosis or occlusion may be appropriate, although these usually cause transitory ischaemic attacks or strokes, rather than a dementia³⁷.

Echocardiography is important if a cardiac source of emboli is considered possible and should be performed in all patients with more than one cortical infarct¹⁸.

Skin biopsy, chromosome and DNA analysis may be helpful in the diagnosis of CADASIL and MELAS. Measuring cerebral

blood flow and metabolism has become increasingly more sophisticated with the advent of positron emission tomography (PET) which uses radioactive tracers to measure metabolism and blood flow. Frackowiak *et al.* showed that cerebral blood flow (CBF) and metabolism fell with increasing dementia in both MID and degenerative dementia³⁸. Normal CBF is 50–70 ml/100 g/min: ischaemia only occurs once the CBF falls below 10–20 ml/100 g/min. Focal abnormalities are found in both MID and degenerative dementia, although in the vascular group the individual focally deranged areas are patchy and match the unique patterns of ischaemic damage, whereas in AD the focal abnormalities are mainly temporoparietal. PET largely remains a research tool.

Single photon emission CT (SPECT) is becoming a more widely available tool, as most X-ray departments have the necessary equipment. Radioactive tracers are used to measure blood flow. Similar temporoparietal abnormalities in AD and patchy irregularities in MD have been reported³⁹. However, all these studies have used a clinical scoring system with/without CT to differentiate MID and AD. None have had pathological confirmation. Because only 10–20% of dementias are multi-infarct, and because an approximate equal percentage have mixed disease (stroke with coincidental AD), it will not be possible to know the true value of PET and SPECT until serial studies are performed with pathological confirmation.

Treatment

As with all vascular disease treatment initially involves dealing with the risk factors—hypertension, hyperlipidaemia, diabetes mellitus and smoking. The Syst-Eur Hypertension trial has demonstrated that treating hypertension prevents dementia⁴⁰. This is combined with specific treatment for the underlying disease process. Aspirin is of proven value in preventing stroke⁴¹. Whether adding dipyridamole to aspirin confers any greater benefit than aspirin alone remains debatable^{42,43}. There is no doubt that clopidogrel is a new effective antiplatelet agent at least as good as aspirin; it should be considered in those intolerant of aspirin⁴⁴.

Warfarin is the treatment of choice in those with atrial fibrillation, but the risk of haemorrhage secondary to warfarin is higher in those with leucoaraiosis^{45,46}. It must be used with caution if poor compliance and falls are a problem. Anti-coagulation should also be considered in the antiphospholipid syndrome¹⁶.

In spite of the many claims, no vasodilator, calcium antagonist or neuroprotective agent has been shown to help vascular dementia. Steroids and immunosuppression may be indicated if an arteritis is proven.

Tatemichi³⁷ has published a case of dementia with bilateral internal carotid occlusions which improved after extracranial–intracranial bypass surgery to improve the blood flow to the brain.

This section has dealt with the specific treatments for vascular disease: it is crucial, of course, to provide symptomatic treatment and a full care package for the many other problems the unfortunate patient with vascular dementia may experience.

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Vascular Dementia Subgroups: Multi-infarct Dementia and Subcortical White Matter Dementia

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Until the 1960s, almost all dementias of old age were considered secondary to “hardening of the arteries” and chronic ischaemia. In the next decades, vascular causes of dementia came to be considered rare, and multi-infarct dementia (MID) almost its only cause. The term “vascular dementia” (VaD) was introduced in the 1990s to emphasize that several cerebrovascular disorders might cause dementia.

An ischaemic stroke increases the risk for dementia several-fold¹. The dementias related to stroke are MID and strategic infarct dementia. MID is related to multiple small or large brain infarcts, often too small individually to produce a major clinical incident. The typical patient has a history of stroke or transitory ischaemic attacks with acute focal neurological symptoms and signs. The clinical picture includes sudden onset, stepwise deterioration and a fluctuating course of the dementia. In the early stages, the cognitive impairment may have a large variability, depending on the site of the lesions. However, in a large minority of cases the dementia may have a gradual onset, with a slowly progressive course² and without focal signs or infarcts on brain imaging (especially when computed tomography has been used), which makes it difficult to differentiate from AD. It has been suggested that the dementia may be related to the location or the volume of the infarcts. The risk factors suggested for MID are similar to those in stroke, including advanced age, male sex, hypertension, diabetes mellitus, smoking and cardiac diseases³.

Ischaemic white matter lesions (WMLs) were first described in cases of dementia by Durand-Fardel in 1854, followed by Binswanger in 1894, and by Alzheimer in 1898. Fewer than 50 autopsied cases were described up to 1980⁴. In the 1980s, when WMLs became possible to discern on brain imaging, they were suddenly reported in thousands of patients. Before the advent of brain imaging, the level of interest in white matter disorders among pathologists was low and the white matter was not routinely evaluated in most patients. The pathologic description

includes marked or diffuse demyelination and moderate loss of axons, with astrogliosis and incomplete infarction in subcortical structures of both hemispheres, and arteriosclerotic changes, with *hyalinization or fibrosis and thickening of the vessel walls and narrowing of the lumen of the small penetrating arteries and arterioles in the white matter*. The main hypothesis regarding the cause of WMLs is that long-standing hypertension causes *lipohyalinosis and thickening of the vessel walls, with narrowing of the lumen of the small perforating arteries and arterioles which nourish the deep white matter*⁴. The dementia is probably caused by subcortical–cortical or cortico–cortical disconnection, and is generally of a subcortical type with extrapyramidal signs, especially psychomotor retardation. A frontal lobe syndrome with apathy, loss of drive and emotional blunting is common.

Regarding the subtypes of VaD, there is a discussion between lumpers and splitters. In the NINDS–AIREN criteria⁶, MID and WMLs are lumped together and it has been questioned whether present scientific knowledge permits anything more than a broad definition of VaD. Focal neurological deficits and brain infarcts are common in subjects with WMLs, and the presence of WMLs are reported to increase the risk for dementia in subjects with stroke. The reason for the common co-occurrence of stroke and WMLs may be that they share similar risk factors, mainly hypertension. However, not all demented subjects with WMLs have cortical infarcts in their brains, and not all individuals with MID have WMLs. Therefore, it may be better to adopt the splitting view to better categorize these entities, with the understanding that there is a large overlap.

There is also an overlap between VaD and AD. WMLs have been described in a high proportion of cases with AD on both brain imaging and at autopsy, and they seem to be more common in late-onset than in early-onset AD⁴. It has even been suggested that white matter degeneration precedes and causes the cortical atrophy in AD⁵. A considerable proportion of subjects from the general population fulfilling the research criteria for AD or VaD

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have mixed pathologies^{7,8}. Cerebrovascular diseases may increase the possibility that individuals with AD lesions in their brains will express a dementia syndrome⁹, and "post-stroke dementia" is often a mixture of the direct consequences of stroke, pre-existing AD pathology and the additive effects of these lesions and ageing¹⁰. The coincidence of AD and VaD may even be the most common form of dementia.

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The Role of Blood Pressure in Dementia

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The association between hypertension and dementia/cognitive impairment has received increased interest recently. Both the Honolulu–Asia Aging Study¹ and the Framingham Study² report that low performance in psychometric tests in the population is related to a higher systolic blood pressure decades before the measurement of cognitive function. We found that systolic and diastolic blood pressure was increased 10–15 years before the onset of both Alzheimer's disease (AD) and vascular dementia, as well as in individuals with white matter lesions³. Furthermore, middle-aged non-demented hypertensive individuals have an increased amount of senile plaques and neurofibrillary tangles, the histopathological hallmarks of AD, in their brains⁴. In cross-sectional studies, blood pressure is generally negatively correlated to cognitive performance before age 75, while there is a positive correlation above that age. Furthermore, elderly subjects with already manifested AD and vascular dementia have lower blood pressure than the non-demented⁵. Thus, although previously high blood pressure seems to be related to cognitive decline and dementia, low blood pressure is often related to already manifested dementia.

Several mechanisms may be involved in the association between hypertension and dementia/cognitive decline. First, the risk of stroke increases with increasing blood pressure, and stroke is a strong risk factor for dementia. It is thus reasonable to believe that high blood pressure is a risk factor for stroke-related dementia. Hypertension is also a risk factor for ischaemic subcortical white-matter lesions (WMLs)⁶, which are often found in subjects with late-onset AD and multi-infarct dementia (MID). The main hypothesis regarding the cause of WMLs is that longstanding hypertension causes lipohyalinosis and thickening of the vessel walls, with narrowing of the lumen of the small perforating arteries and arterioles that nourish the deep white matter. Episodes of hypotension may lead to hypoperfusion and hypoxia–ischaemia, leading to loss of myelin in the white matter. It has been suggested that the arterial changes are due to exposure

of vessel walls to increased pressure over time. The greater the pressure and/or lifespan, the more likely are these changes to be present.

A third explanation is that chronic hypertension may lead to a dysfunction of the blood–brain barrier (BBB), with increased permeability of the endothelial cell layer and extravasation of serum proteins⁷. The CSF:serum albumin ratio is a generally accepted method of assessing the BBB function in living subjects. We recently found that AD, MID and severe forms of WMLs were all associated with an increased CSF:serum albumin ratio in a population-based study of 85 year-olds⁸. A breakdown in the BBB may cause brain lesions by cerebral oedema, activation of astrocytes, or destructive enzymes or other poisons that pass through the damaged vessel walls. Fourth, the renin–angiotensin system is an example of a system that may be involved in the pathogenesis of both hypertension and dementia⁹. Its effector peptide, angiotensin II, has several blood pressure increasing effects, promotes hyperplasia and hypertrophy in vascular smooth muscle cells, and affects memory and behaviour. Fifth, psychological stress has been suggested to be a risk factor for both hypertension and dementia¹⁰.

The association between low blood pressure and dementia has several explanations. First, systemic hypotension associated with reduced cerebral blood flow may give rise to a spectrum of ischaemic neuronal lesions in the brain and may also lead to ischaemic loss of myelin in the white matter. Second, several of the blood pressure-regulating areas in the central nervous system are affected in dementia disorders. Therefore, dementia disorders and their associated brain changes may *per se* influence the blood pressure⁵. A correlation between the number of C-1 neurons in the medulla oblongata and blood pressure has been reported in AD¹¹. Blood pressure decline during the course of AD, and low blood pressure was related to cerebral atrophy in non-demented 85 year-olds⁵. It thus seems likely that low blood pressure in demented individuals is secondary to the brain lesions. If cerebral disorder

have mixed pathologies^{7,8}. Cerebrovascular diseases may increase the possibility that individuals with AD lesions in their brains will express a dementia syndrome⁹, and “post-stroke dementia” is often a mixture of the direct consequences of stroke, pre-existing AD pathology and the additive effects of these lesions and ageing¹⁰. The coincidence of AD and VaD may even be the most common form of dementia.

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The Role of Blood Pressure in Dementia

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The association between hypertension and dementia/cognitive impairment has received increased interest recently. Both the Honolulu–Asia Aging Study¹ and the Framingham Study² report that low performance in psychometric tests in the population is related to a higher systolic blood pressure decades before the measurement of cognitive function. We found that systolic and diastolic blood pressure was increased 10–15 years before the onset of both Alzheimer’s disease (AD) and vascular dementia, as well as in individuals with white matter lesions³. Furthermore, middle-aged non-demented hypertensive individuals have an increased amount of senile plaques and neurofibrillary tangles, the histopathological hallmarks of AD, in their brains⁴. In cross-sectional studies, blood pressure is generally negatively correlated to cognitive performance before age 75, while there is a positive correlation above that age. Furthermore, elderly subjects with already manifested AD and vascular dementia have lower blood pressure than the non-demented⁵. Thus, although previously high blood pressure seems to be related to cognitive decline and dementia, low blood pressure is often related to already manifested dementia.

Several mechanisms may be involved in the association between hypertension and dementia/cognitive decline. First, the risk of stroke increases with increasing blood pressure, and stroke is a strong risk factor for dementia. It is thus reasonable to believe that high blood pressure is a risk factor for stroke-related dementia. Hypertension is also a risk factor for ischaemic subcortical white-matter lesions (WMLs)⁶, which are often found in subjects with late-onset AD and multi-infarct dementia (MID). The main hypothesis regarding the cause of WMLs is that longstanding hypertension causes lipohyalinosis and thickening of the vessel walls, with narrowing of the lumen of the small perforating arteries and arterioles that nourish the deep white matter. Episodes of hypotension may lead to hypoperfusion and hypoxia–ischaemia, leading to loss of myelin in the white matter. It has been suggested that the arterial changes are due to exposure

of vessel walls to increased pressure over time. The greater the pressure and/or lifespan, the more likely are these changes to be present.

A third explanation is that chronic hypertension may lead to a dysfunction of the blood–brain barrier (BBB), with increased permeability of the endothelial cell layer and extravasation of serum proteins⁷. The CSF:serum albumin ratio is a generally accepted method of assessing the BBB function in living subjects. We recently found that AD, MID and severe forms of WMLs were all associated with an increased CSF:serum albumin ratio in a population-based study of 85 year-olds⁸. A breakdown in the BBB may cause brain lesions by cerebral oedema, activation of astrocytes, or destructive enzymes or other poisons that pass through the damaged vessel walls. Fourth, the renin–angiotensin system is an example of a system that may be involved in the pathogenesis of both hypertension and dementia⁹. Its effector peptide, angiotensin II, has several blood pressure increasing effects, promotes hyperplasia and hypertrophy in vascular smooth muscle cells, and affects memory and behaviour. Fifth, psychological stress has been suggested to be a risk factor for both hypertension and dementia¹⁰.

The association between low blood pressure and dementia has several explanations. First, systemic hypotension associated with reduced cerebral blood flow may give rise to a spectrum of ischaemic neuronal lesions in the brain and may also lead to ischaemic loss of myelin in the white matter. Second, several of the blood pressure-regulating areas in the central nervous system are affected in dementia disorders. Therefore, dementia disorders and their associated brain changes may *per se* influence the blood pressure⁵. A correlation between the number of C-1 neurons in the medulla oblongata and blood pressure has been reported in AD¹¹. Blood pressure decline during the course of AD, and low blood pressure was related to cerebral atrophy in non-demented 85 year-olds⁵. It thus seems likely that low blood pressure in demented individuals is secondary to the brain lesions. If cerebral disorder

causes low blood pressure, the question arises whether cerebral changes may induce high blood pressure. Recently, it was shown that infusion of $A\beta_{42}$ (a protein deposited in the brains and cerebral vessels of AD victims) increased blood pressure in anaesthetized rats, suggesting that circulating levels of this protein may exert vasopressor actions *in vivo*¹².

Treatment of hypertension may thus have a preventive effect on cognitive decline and dementia. The recent finding from the Syst-Eur trial¹³, that treatment of isolated systolic hypertension reduces the incidence of dementia by 50%, supports this hypothesis, but the number of demented in that study was small. It has been suggested that overtreatment with antihypertensive drugs may increase the risk of dementia in the very old by causing cerebral hypoperfusion. No studies so far provide support for this opinion. We recently reported¹⁴ that subjects who became demented during a 15-year follow-up used antihypertensive drugs less often than those who did not become demented. Although these findings do not preclude the possibility that overtreatment of hypertension may cause ischaemia in the brain in a subset of individuals, they do not support the hypothesis that antihypertensive treatment may cause cognitive impairment in the elderly.

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Neuropathology: Other Dementias

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There are many causes of dementia in addition to Alzheimer's disease (AD) and vascular dementia, and the commonest of these seems to be dementia with Lewy bodies (DLB), which perhaps accounts for 10% of all cases of dementia, although the precise incidence depends on the population studied and the pathological criteria used for diagnosis^{1,2}. In a hospital series, it may be second only to AD as a cause of dementia³. After DLB, other causes are distinctly rare and include Creutzfeldt–Jakob disease, Huntington's disease and the frontotemporal dementias.

DEMENTIA IN PARKINSON'S DISEASE AND DEMENTIA WITH LEWY BODIES

Parkinson's disease is a disorder characterized by rigidity, tremor and bradykinesia, usually with an onset in the sixth decade. Neuronal loss and gliosis are typically evident in the substantia nigra and loci coerulei, but also in the dorsal vagal nuclei, dorsal raphe nuclei and nucleus basalis of Meynert. Lewy bodies, spherical intracytoplasmic inclusions composed of granular and filamentous elements, immunoreactive with antibodies against neurofilament protein, paired helical filaments, ubiquitin and, in aminergic neurones, tyrosine hydroxylase protein^{4–7}, are found not only in pigmented brainstem neurones but also in over 20 different nuclei, pigmented and non-pigmented⁸. Changes in aminergic pathways, particularly dopaminergic, are well documented⁴.

Dementia has been reported to affect 34.6% of patients with parkinsonism in one large autopsy series and it seems likely that a number of different mechanisms are responsible⁹.

Concurrent AD

It has been claimed that AD is six times more common in patients with Parkinson's disease than in age-matched controls¹⁰ and 12–25% of demented parkinsonian patients in one series had AD⁴.

In Jellinger's study⁹, most of the demented parkinsonians also had the pathological changes of AD. Other studies have shown that cortical senile plaques and neurofibrillary tangles are more numerous in demented patients with Parkinson's disease than non-demented Parkinson's disease sufferers or controls^{11–13}. The wide spectrum of dementia in Parkinson's disease has been attributed to the overlap of clinical and subclinical pathological changes in both diseases¹⁴.

Innominato-cortical Dysfunction

Neuronal loss from the basal nucleus of Meynert (BNM) is greater in demented than in non-demented parkinsonian patients¹¹ and this depletion may be 40–80% in the former as opposed to 20–50% in the latter^{15,16}.

Furthermore, there is also evidence of an accompanying reduction in choline acetyltransferase (CAT) and acetylcholinesterase in the BNM and all areas of the cerebral cortex. Decreased CAT activity in the temporal cortex correlates with BNM neuronal loss in Parkinson's disease, unlike AD, and also correlates with memory impairment, although not with the numbers of cortical plaques and tangles¹⁷.

Neuronal Loss in Pigmented Brainstem Nuclei

It is generally stated that there is no correlation between neuronal loss in the substantia nigra and mental impairment¹¹, but it has been found that neuronal loss in the medial substantia nigra correlates with the severity of dementia, presumably due to loss of projections to the caudate nuclei, limbic system and cerebral cortex¹⁸. In addition, severe neuronal loss and subnormal noradrenaline metabolism in the loci coerulei are commoner in demented than non-demented parkinsonian patients^{11,19}. Medial substantia nigra loss, together with cortical AD changes, seemed to underlie dementia in one study, whereas depression in Parkinson's disease was associated with more severe loss of neurones from the dorsal raphe nuclei¹³.

Dementia with Lewy Bodies

Small numbers of Lewy bodies may be present in the cerebral cortex in patients without neurological impairment, or with Parkinson's disease. However, a disorder characterized by early neuropsychiatric features, dementia, visual hallucinations, fluctuating conscious level and relatively mild Parkinson's disease justifies separate identification as DLB²⁰ because of different therapeutic implications. Many cases also have large numbers of brainstem Lewy bodies and it seems likely that DLB, with a predominantly cortical distribution of Lewy bodies, lies at one end of a spectrum of Lewy body disorders, in which idiopathic Parkinson's disease, with Lewy bodies predominantly localized in the brainstem, is the opposite end²¹.

In many reported cases, cortical Lewy bodies have been associated with AD-type changes, leading to the conclusion that

the cause of dementia in diffuse LBD is cortical^{22,23}. However, the distribution of neuritic plaques and neurofibrillary tangles does not necessarily parallel the distribution of Lewy bodies²⁴ and, in some cases with Lewy bodies in the cerebral cortex and brainstem nuclei, AD-type changes are not present²⁵.

The precise nosological relationship between DLB and AD, and between DLB and patients with Parkinson's disease who subsequently become demented, requires elucidation³.

CREUTZFELDT–JAKOB DISEASE

Creutzfeldt–Jakob disease (CJD) belongs to the family of transmissible spongiform encephalopathies affecting humans and animals²⁶. In humans the main forms are sporadic CJD, iatrogenic CJD (associated mostly with pituitary hormones prepared from cadaver pituitaries), familial forms, including Gerstmann–Straussler–Scheinker (GSS) syndrome and variant CJD (vCJD).

Sporadic CJD is typically a rapidly progressive dementia with associated myoclonus, but a number of different clinical and pathological variants have been described^{27,28}. The worldwide incidence is approximately 1–2/million/year²⁹, although the average annual incidence in England and Wales was 0.3 in 1970–1979³⁰, and 0.49 in 1980–1984³¹. The highest reported incidence of 75 is amongst Libyan Jews³². Although often a presenile dementia, the peak age-specific mortality rate in England and Wales is in the seventh decade, with a mean of 63.2 years and a range of 33–82³¹.

Approximately 10–15% of cases of CJD are familial and seem to follow an autosomal dominant pattern of inheritance³³. GSS is also familial, but clinically and pathologically more closely resembles kuru than CJD, although caused by the same transmissible agent^{34,35}.

To date, there has been only one reported case of vCJD in the elderly^{36a}. All of the other cases have been younger (mean age at death 29 years). The presenting features in both age groups are behavioural change and sensory symptoms with later development of cerebellar ataxia, dementia and myoclonus³⁶.

Although pathological variants of CJD have been described, the principal features are fluctuating spongiform change of the grey matter, with neuronal loss, and astrocytic hypertrophy and hyperplasia³⁷. Amyloid plaques, particularly in the cerebellum, are present in 5–10% of cases of CJD and the majority of cases of kuru and GSS³⁴. Florid amyloid plaques of prion protein are also characteristic of vCJD.

The transmissible nature of CJD has been known for many years^{37,38}. Prions^{39,40} as proteinaceous infectious particles consisting of one protein (PrP 27-30), which is encoded by a cellular gene and derived from a larger protein. PrP 27-30 is inseparable from the infectivity of the scrapie agent and constitutes the amyloid of cerebral plaques in “prion” diseases. The PrP or a closely-related gene clearly controls the clinical expression of the disease.

The causal relationship of PrP with the disease is strengthened by the presence of disease-specific mutations linked to, or tracking with, inherited “prion” diseases, such as the substitution of leucine for proline at PrP codon 102 in ataxic forms of GSS⁴¹ and the substitution of lysine for glutamate at codon 200 in Libyan Jews⁴². Similarly, the valine 129 homozygous genotype is associated with susceptibility to iatrogenic CJD⁴³ and methionine 129 homozygosity with vCJD³⁶.

The continued unravelling of the molecular biology of the prion diseases is likely to reveal as many surprises in the future as have already been provided by this fascinating group of

disorders. However, it seems improbable that “prion dementias” constitute a large group of as yet unrecognized neurodegenerative disorders and, for all practical purposes, transmissible spongiform encephalopathies in humans and “prion dementias”, are the same⁴⁴.

HUNTINGTON'S DISEASE

Huntington's disease (HD) is a disorder of midlife onset (mean age 41 years) characterized by progressive chorea, psychological changes and dementia, although not necessarily in that order. The disease is encountered in the senium, sometimes because of prolonged survival of presenile cases, but also because chorea only appears in 28% of HD patients after the age of 50 and, occasionally, not until the seventh or eighth decade⁴⁵. HD is inherited as an autosomal dominant trait with 100% penetrance and the HD gene has been localized near the telomere on the short arm of chromosome 4⁴⁶.

The principal neuropathological features are neuronal loss and gliosis in the corpus striatum, and a five-grade system has been devised for pathologically classifying HD⁴⁷, which closely correlates with the degree of clinical disability⁴⁸. Most of the neuropathologically studied cases fall into Grades 2 and 3 (79%)⁴⁹; 50% of neurones have been lost from the caudate nucleus by the time grade 1 pathological changes are recognizable and 95% by grade 4. There is an accompanying increase in astrocytes and oligodendrocytes, reaching a maximum that is significantly different from control material in grade 4 for the former and grades 0–2 for the latter⁴⁷.

The distribution of neuronal loss within the corpus striatum is non-uniform. In Grades 1 and 2 the medial caudate nucleus is more severely affected than the lateral, although this distinction is lost when the pathological changes become more severe⁴⁷. The anteroventral part of the putamen is also spared⁵⁰. Not only is the disease process in the corpus striatum anatomically variable, but also not all neuronal populations are equally affected. The small to medium-sized spiny neurones (which have a large synaptic surface and distant connections) are most affected in HD, and these neurones contain γ -aminobutyric acid (GABA) enkephalins and substance P^{51,52}. On the other hand, medium aspiny neurones, containing NADPH-diaphorase, somatostatin and neuropeptide Y, and large cholinergic neurones, are relatively spared^{53,54}. The decreased concentration of GABA and enzymes involved in its metabolism, which has been known for many years⁵⁵, together with the decreased concentrations of substance P and the enkephalins, can be explained on the basis of selective loss of striatal neuronal populations.

The disease is caused by expansion of a trinucleotide CAG repeat beyond 35 repeats in the first exon of the huntingtin gene on chromosome 4⁵⁶. The CAG repeats are translated to polyglutamine repeats in huntingtin and, although the normal function of huntingtin is incompletely understood, it appears to be associated with the cytoskeleton and necessary for neurogenesis, and expression of mutant huntingtin is necessary for disease development. The ability of huntingtin to interact with other proteins is influenced by CAG length and it is possible that neuronal apoptosis may be triggered by abnormal interaction with caspase 3^{57,58}. Furthermore, there is also evidence of abnormal metabolism in HD brain tissue, resulting in the generation of free radicals, so that more than one mechanism may be operating to produce cell death in HD⁵⁹.

Although corpus striatum changes dominate the pathological picture in HD, other areas of the brain are involved. The

cerebral cortex showed 21–29% loss of substance, the cerebral white matter 29–34% and the thalamus 28% in one study⁶⁰, but cortical abnormalities may be very difficult to identify⁴⁹. Neuronal loss has been identified in the superior frontal and cingulate cortices, particularly of laminae III and V pyramidal neurones, but this does not seem to correlate with severity of striatal pathology, suggesting that it is a primary part of the disease process⁶¹.

However, another study⁶² found evidence of loss of large pyramidal neurones from laminae III, V and VI of the prefrontal cortex, which was most severe in pathological grade 4 HD, and suggested that this represented retrograde degeneration of cortical glutamatergic neurones. Nerve cell loss from specific regions of the entorhinal cortex and subiculum has also been reported⁶³.

There is controversy over whether neuronal loss in the substantia nigra does⁶⁴ or does not⁶⁵ occur in HD. Preservation of neuronal populations in the nucleus basalis of Meynert, locus coeruleus and dorsal raphe nuclei has also been reported, leading to the suggestion that dementia in HD may be due not to damage to subcortical nuclei, as in the other “subcortical” dementias, but to failure of ascending systems within the basal ganglia or neuronal loss in the cerebral cortex⁶⁵.

FRONTOTEMPORAL DEMENTIA

The frontotemporal dementias (FTD) are non-Alzheimer forms of dementia characterized clinically by behavioural and personality change leading to apathy and mutism, and by progressive atrophy of the frontal, anterior parietal and anterior temporal lobes. The histology is variable and defines three separate subgroups⁶⁶:

- *Frontal lobe degeneration*. This is characterized by microvacuolar degeneration, gliosis and neuronal loss, principally affecting laminae II and III.
- *Pick's disease* (FTD with Pick-type histology) is represented by transcortical neuronal loss, gliosis and cavitation, together with the presence of tau and ubiquitin-positive intraneuronal inclusions (Pick bodies) and α B crystallin-positive ballooned neurones (Pick cells). Although there is overlap between Pick's disease and other forms of FTD, as well as with corticobasal degeneration, the distinctive distribution of the pathological changes—such as the involvement of the dentate fascia by Pick bodies—has led some authors to regard Pick's disease as a unique disease rather than a histological variant within the FTD spectrum⁶⁷. However, both of these different histologies can have a variable topographical distribution in the brain, producing progressive language disorder when both temporal lobes are involved or the disease primarily affects the left hemisphere, or progressive apraxia when parietal and motor areas are involved⁶⁶.
- *Motor neurone disease inclusion dementia*. Either of these histologies can overlap with classical motor neurone disease, either as a dementing disorder occurring in patients known to have MND, or also as an identical histology occurring in patients without motor dysfunction^{66,68}.

Fifty-eight percent of cases of FTD show a previous family history of a similar disorder, and linkage to chromosome 17 has been established in some families. Following the current trend for “molecular” classification of the dementias, the various forms of FTD have sometimes been described as “tauopathies” but 64% cases of FTD in one study showed no intracellular tau-positive inclusions⁶⁹. This is an area where further study and clarification is required.

OTHER DISORDERS

Many other disorders may be associated with dementia in the elderly, including progressive supranuclear palsy⁷⁰ and, occasionally, multiple sclerosis⁷¹. There are also a number of rare dementing disorders of uncertain nosological status, such as the tangle-predominant form of dementia⁷² and argyrophilic grain disease⁷³.

This section of the book would not be complete without the reader being reminded of the estimated 13.2%⁷² of causes of dementia in the elderly—including intracranial space-occupying lesions, particularly chronic subdural haematoma; metabolic disorders, such as hypothyroidism and hypopituitarism⁷⁵; and “normal-pressure” hydrocephalus—that are potentially treatable⁷⁶.

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Dementia and Parkinson's Disease

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Although James Parkinson, in his 1817 monograph, believed that this disease left "the senses and intellects . . . uninjured", there has been increasing realization that Parkinson's disease (PD) is associated in many cases with dementia. The dementia has specific clinical and psychological characteristics and is now believed to be associated in the majority with specific neuropathological changes. Certainly, dementia in elderly patients with PD complicates their management, limits treatment and survival and leads to a heavy burden on medical and welfare services and family alike. The dementia of PD is therefore a subject of great importance, in which substantial advances in understanding have taken place in recent years.

In cases where parkinsonism and dementia coexist, the differential diagnosis must include Lewy body dementia (LBD), progressive supranuclear palsy, corticobasal degeneration, meso-limbo-cortical dementia and severe Alzheimer's disease. In life, LBD is characterized by the association of parkinsonism and a dementia with deficits in attention, executive functions and visuospatial abilities. There are often fluctuations in alertness, sometimes amounting to transient unresponsiveness. And recurrent formed hallucinations with little or no affective component are common. In spite of these typical features, the distinction of LBD in the elderly may be very uncertain.

About 20% of all patients with PD develop dementia. The prevalence of diagnosed PD is overall about 1/1000, but the prevalence increases with age and reaches nearly 2% in the ninth decade. An undiagnosed pre-symptomatic PD is very much more common, so that pathological findings indicative of the disease are present in about 10% of the healthy population in the ninth decade³. Dementia in PD is more common in the elderly patient, and patients with dementia tend to be significantly older at the onset of their PD⁴. By contrast, early-onset PD tends to have a low incidence of dementia⁵.

Patients who present with dementia and later develop typical PD are probably suffering from LBD, now categorized as one of the "Lewy body disorders".

Dementia associated with PD is therefore a common occurrence in clinical practice. A number of important and interesting questions arise from these facts. Has the dementia any specific clinical features that distinguish it from "senile" dementia of the Alzheimer type? Has the type of PD that is associated (perhaps at a late stage) with dementia any specific clinical features compared with the type of disease where dementia is uncommon? What is the relationship, if any, between treatment for PD and the development of dementia? What are the neuropathological correlates between PD and dementia? Are both phenomena caused by the same process?

CLINICAL FEATURES OF THE DEMENTIA ASSOCIATED WITH PARKINSON'S DISEASE

Cognitive testing in PD may be affected by a number of extraneous factors, including impaired motor function, medication or mood disturbance. In people with no associated disease, cognitive function can be compared with premorbid intelligence and the decline estimated. However, no validated method for estimating premorbid intelligence in PD has been developed. Depression is common in PD and may give a pseudodementia, relieved by antidepressant treatment. For these and other reasons, no generally accepted pattern of cognitive function has emerged as characteristic of PD and longitudinal studies are needed.

ORGANIC DISORDERS

The concept of subcortical dementia was originally introduced by Albert with reference to the dementia in the rare form of parkinsonism seen in progressive supranuclear palsy⁷. Subcortical dementia is characterized by forgetfulness, slowing of thought processes (bradyphrenia), apathy and impaired ability to manipulate acquired knowledge. This pattern of deficit is contrasted with diseases such as Alzheimer's disease, characterized by global memory impairment, aphasia, agnosia and apraxia. While much overlap seems to exist between these types of cognitive impairment, bradyphrenia, apathy and depression seem to characterize the dementia of PD. Thought-block is an early characteristic of the dementia and fluctuation of the cognitive impairment has been noted⁸. Visual hallucinations, often with little or no affective component, are common. Less common are paranoid delusional states.

Patients with PD and dementia show a treatment response for their parkinsonism that is inversely related to the severity of the cognitive changes⁹.

DO SPECIFIC CLINICAL FEATURES OF PARKINSON'S DISEASE INDICATE THE LIKELIHOOD OF DEMENTIA?

Lieberman *et al.*¹⁰ demonstrated that the dementia of PD was associated with a later age of onset, a poorer response to L-dopa and less abnormal involuntary movements and dose fluctuations in response to L-dopa. These findings have been confirmed^{4,5,9} in later studies.

Tremor is less common as a presenting symptom in patients with PD associated with dementia, whereas dyspraxia of gait and

dyspraxias for hand movements are more common. Thus, a patient who has difficulty in performing the discrete finger movements used to test for bradykinesia may commonly show early cognitive changes.

The rate of progression of the PD is significantly greater in cases where dementia further affects management and survival. Patients with dementia tolerate drug treatment less well and tend to develop more persistent and severe toxic confusional states.

Thus, many authors accept that PD may exist in a late-onset form with certain characteristic clinical features and a relatively poor response to L-dopa where dementia is common. By contrast, there is a type of PD in which dementia is rare. The disease has an earlier onset and response to L-dopa is initially very good^{12,13}.

All drugs used in the treatment of PD carry a risk of provoking toxic confusional states and this reaction is more likely and more severe in patients with cognitive changes. Where therapeutic benefit from a drug is likely to be modest, there is a relative contraindication to the use of that drug in an elderly patient, particularly if they are showing cognitive changes. For this reason, the anticholinergic drugs are seldom recommended in geriatric practice¹⁴.

Conversely, in patients with classical PD in middle life, where the prospect of cognitive impairment is small (but the risks of permanent response fluctuations to L-dopa are high) anticholinergic drugs are indicated in preference to L-dopa if disability can be satisfactorily controlled.

The fluctuation of the dementia already noted may suggest that the cognitive changes are related to drug therapy. This possibility should usually be investigated by dose reduction or withdrawal, but if no improvement occurs the patient need not be deprived of the physical benefits of the drug. Many elderly patients with mild cognitive changes tolerate L-dopa therapy with benefit and without any significant exacerbation of their mental impairment. Occasionally the administration of L-dopa in patients with dementia may provoke states of excitement with impulsive behaviour and aggravation of the dementia¹⁵.

The synthetic dopamine agonists (bromocriptine, lisuride, pergolide, ropinirole) and the anticholinergics should be used only with great caution in the parkinsonian patient with dementia.

WHAT IS THE RELATIONSHIP, IF ANY, BETWEEN TREATMENT FOR PARKINSON'S DISEASE AND THE DEVELOPMENT OF DEMENTIA?

The two drugs in most widespread use for the treatment of PD are benzhexol and L-dopa (with decarboxylase inhibitor). We must examine the evidence concerning whether either of these drugs could be causing or accelerating dementia in PD.

Alzheimer's disease is known to be associated with a reduction in brain choline acetyltransferase, and anticholinergic drugs exacerbate the neural changes in this condition, but there is no evidence of permanent effects on higher mental function from these drugs in PD. Confusional states provoked by benzhexol recover with a timescale of usually less than 4 weeks. It is noteworthy that the use of a high-dose anticholinergic drug in the treatment of dystonias over many years is not associated with any cognitive changes.

However, studies of the prevalence of dementia in PD before the L-dopa era gave figures of 3.2–10%¹⁶ or 851%¹⁷, whereas since the introduction of L-dopa, the prevalence of dementia has risen to above 20%.

Like anticholinergics, L-dopa may provoke acute confusional states that recover when the drug is withdrawn. But withdrawal of L-dopa rarely alters the decline of mental faculties in parkinsonian patients with dementia. The duration of L-dopa treatment does not appear to correlate with the likelihood of mental

impairment¹⁸. At the present time, the evidence is against L-dopa treatment causing or accelerating the dementia of PD.

WHAT ARE THE NEUROPATHOLOGICAL CORRELATES BETWEEN PARKINSON'S DISEASE AND DEMENTIA?

With the advent of staining techniques based on anti-ubiquitin immunocytochemistry¹⁹ neuropathological correlates between PD and dementia showed that by far the commonest cause of dementia in this condition was diffuse LBD²⁰. Previously parkinsonian dementia had been attributed to a supposed association between PD and Alzheimer's disease, but the new staining techniques showed that in PD without dementia the Lewy body was substantially confined to the brainstem, whereas in PD with dementia, diffuse cortical Lewy body formation was demonstrable.

By contrast, in Alzheimer's disease, neuronal loss in the nucleus basalis is associated with neurofibrillary tangles in this region and in the dorsal raphe nucleus. In cases of PD with dementia, scanty tangle formation may be seen in the hippocampus or neocortex, but with profuse cortical Lewy bodies not seen in Alzheimer's disease.

Senile plaques have been recommended as an important neuropathological criterion for the diagnosis of Alzheimer's disease²³. But senile plaques are commonly seen in similar density to cortical Lewy bodies in cases with dementia²⁰, and thus fail to distinguish between the two conditions.

The dementia seen in some cases of PD is attributable to neuronal cell loss in the cortex associated with diffuse cortical Lewy body formation.

ARE BOTH PHENOMENA CAUSED BY THE SAME PROCESS?

Lewy body disease, in which parkinsonism is associated with dementia, is probably the second commonest cause of dementia after Alzheimer's disease²⁴. The Lewy body is an intra-neuronal eosinophilic inclusion body that shows ubiquitin immunoreactivity, and is the pathological hallmark of PD.

Thus, PD is characterized by Lewy body formation in the brainstem and particularly in the substantia nigra²⁵. In demented cases of PD there is cortical Lewy body formation as well and the severity of the dementia correlates with cortical Lewy body density²⁰. It is therefore reasonable to conclude that the dementia and the parkinsonism are due to the same pathological process. The nature of this process is currently the subject of intense research.

LBD studied post mortem may show a variety of patterns. In the typical case, Lewy bodies may be distributed in the brainstem, diencephalon, limbic system and cortex, but in some cases Lewy bodies are confined to the brainstem and are few or absent in the cortex or limbic system. In these cases there are also no senile plaques, whereas where Lewy bodies are present in the cortex, senile plaques may also be seen.

Probably the most specific marker for the Lewy body is α -synuclein, and this protein has been associated in some families with autosomal dominant PD. The pathological process of Lewy body diseases may be partly genetic and seems to be independent from, but sometimes co-existent with, the pathological process of Alzheimer's disease. There is increasing interest in the genetic markers of PD and dementia, but it remains likely that there is an environmental trigger in genetically susceptible cases.

It has already been noted that parkinsonism with dementia is a form of PD that tends to have later onset than classical PD

without dementia. This suggests that the elderly cerebral cortex is vulnerable to the same pathological processes as the substantia nigra (and which lead to parkinsonism), whereas the younger cerebral cortex is relatively resistant, so that early-onset PD tends to be unassociated with dementia. The nature of this age-related vulnerability of the cortex is unknown, but it is of interest that age-related differential vulnerability to 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) has been demonstrated in primates²⁶.

CONCLUSIONS

The dementia of PD is entirely distinct from the dementia of Alzheimer's disease. In PD dementia is associated with Lewy body formation in the cerebral cortex as well as in the substantia nigra; whereas in Alzheimer's disease neuronal loss and neurofibrillary tangle formation is characteristically in the dorsal raphe nuclei and nucleus basalis of Meynert.

The clinical features of the two forms of dementia are usually distinguishable. More difficult to distinguish clinically, especially in the earlier stages, are the two types of PD. In one the disease presents typically in middle life, progresses slowly, responds well to dopamine agonists and is not associated with dementia; whereas the other (diffuse Lewy body type) presents in late life, progresses more rapidly, responds less well to dopamine agonists and is associated with dementia. The clinical significance of making this distinction is to define prognosis and also to affect management, in particular the use of dopamine agonists and the management of dementia.

The cause of PD and of Alzheimer's disease remains unknown but there are promising lines of investigation related to the genetics and molecular biology of the Lewy body and the neurofibrillary tangle, respectively.

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Clinical Criteria for Dementia with Lewy Bodies

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Why are diagnostic criteria for dementia with Lewy body (DLB) important? There are many reasons, but the most significant are to properly advise patients and carers, to minimize neuroleptic prescribing and possibly to effectively target cholinesterase inhibitor use. Also, since DLB is relatively common, it needs to be routinely excluded in the differential diagnosis of other dementia subtypes, particularly when a diagnosis of Alzheimer's disease (AD) is being considered. Several studies have now reported on either the sensitivity (proportion of cases positively identified) or specificity (proportion of negative cases correctly identified) of the International Consensus criteria for DLB, against neuropathological diagnosis. Most find "probable DLB" criteria to have a specificity 40.8, a figure comparable with the best clinical criteria for AD and Parkinson's disease (PD). High specificity means that when a diagnosis of probable DLB is made, it is likely to be correct. The more lenient "possible DLB" category should be useful as a screening tool for identifying cases in the clinic, although many false-positive diagnoses will be made.

Sensitivity rates for probable DLB are more variable and generally lower. This may in part be due to *retrospective* application of the criteria to case records in most of these studies. Spontaneous documentation of fluctuation and detailed psychiatric phenomenology in case notes is notoriously incomplete, leading some to conclude that inter-rater reliability for individual diagnostic items, especially "fluctuation", is unsatisfactory. Although a tighter operational definition, or a biological measure, of fluctuation would undoubtedly be useful, studies *prospectively* applying the DLB diagnostic criteria generally find inter-rater

reliability for individual items (including fluctuation) and for a final diagnosis of DLB, to be acceptable (k40.6), allowing for diagnostic sensitivities 40.8 to be achieved.

Ancillary investigations will have an important future role in improving the accuracy of clinical diagnosis of DLB. FP-CIT SPECT brain imaging demonstrates a large reduction in dopamine transport to the striatum in DLB, and this may be apparent before extrapyramidal signs are manifest. Nigrostriatal dopaminergic depletion is only rarely seen in AD or vascular dementia. Relative lack of medial temporal lobe atrophy on CT/MRI is also characteristic of DLB compared to AD.

The Second International Workshop on DLB recommended that the Consensus criteria should continue to be used in their current format for recruiting cohorts of DLB patients for research studies and clinical trials. Depression and REM sleep behaviour disorder were suggested as two additional features supporting the diagnosis.

Accurate case detection will ultimately be best achieved by increasing the index of suspicion for a diagnosis of DLB, not only in dementia assessment clinics but also in any setting where elderly patients may present with delirium, movement disorder, falls or syncope. Perhaps we might do better to frame the next revision of the diagnostic criteria within a broader spectrum of Lewy body-related disorders or α -synucleinopathies, thereby acknowledging the links between PD, DLB and primary autonomic failure, and breaking down some currently unhelpful boundaries between psychiatry, neurology and geriatric medicine.

Subcortical Dementia

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Health-care workers who evaluate older patients are frequently confronted with individuals who they suspect have dementia. Although it is often straightforward to determine whether an individual is demented, determining the etiology of the dementia can be difficult. Dementia is not a single disease, but a heterogeneous complex of disorders with the common finding of impairment in multiple cognitive domains. Fortunately, clues leading to a correct diagnosis can be obtained in the physical and mental status examination.

A heuristic device that can aid in the evaluation of patients with dementia is to divide diseases into two categories, based on whether the pathology is primarily cortical or subcortical¹. Although the cortical/subcortical dichotomy has been questioned on pathologic grounds, there are clinical features that distinguish one group from the other. The prototypical cortical dementia is Alzheimer's disease (AD). Diseases that cause subcortical dementia include small vessel cerebrovascular disease, Parkinson's disease and Huntington's disease (Table 50b.1)².

CLINICAL FEATURES OF SUBCORTICAL DEMENTIA

The cardinal clinical features of subcortical dementia are apathy, inattention and psychomotor slowing^{1,3}. Patients with subcortical dementia typically appear apathetic. They have a blunted affect, poor personal hygiene and sloppy appearance. The patients are often aware of their cognitive deficit but appear unconcerned. In contrast, patients in the early stages of AD typically have normal affect, grooming and dress. Patients with AD who are aware of their deficit are typically quite concerned.

It is easy to become frustrated when interviewing a patient with subcortical dementia because the patient's inattention leads to frequent repetition of questions by the examiner. Surprisingly, sometimes minutes after a question is asked, patients with a subcortical dementia will respond with the correct answer. Patients with AD have normal attention.

Slowing is a key feature of subcortical dementia and one of the most obvious signs during the examination. Slowing occurs in multiple areas, including cognition (thought formulation, language generation and processing), sensation (processing of stimuli) and motor performance (bradykinesia). While forcing a patient to respond quickly impairs performance in both cortical and subcortical dementia, individuals with subcortical dementia benefit from being given more time to complete a task. Additional response time does not have a similar beneficial effect for patients with AD. The apathy, inattention and psychomotor slowing due to subcortical dementia can be difficult to distinguish from

Table 50b.1. Causes of subcortical dementia

Parkinson's disease	Small vessel cerebrovascular disease
Progressive supranuclear palsy	Spinocerebellar ataxias
Huntington's disease	Hydrocephalus
Multiple systems atrophy	Multiple sclerosis
Wilson's disease	HIV dementia
Frontal lobe dementia	Vasculitis

depression. Patients with AD do not demonstrate psychomotor slowing, unless they are also depressed.

The type of memory dysfunction differs between subcortical and cortical dementia. Impaired registration and retrieval characterize the memory deficit in subcortical dementia, while intact registration and rapid forgetting are seen in AD. To test registration and retrieval, a patient is asked to repeat a short list of words immediately after presentation (registration) and after a delay (retrieval). In subcortical dementia, the list may have to be presented several times before all items are registered. Registration is intact in AD patients. Patients with subcortical dementia and AD may retrieve a similar number of items, but if presented with a choice, the patient with subcortical dementia can distinguish between items that were presented and items that were not (recognition). AD patients are unable to distinguish between list and non-list items. Patients with subcortical dementia also benefit from hints to recall items (cueing), while AD patients do not.

Patients with subcortical dementia also frequently display deficits in multi-step tasks. They may perform each step of a task individually, but be unable to incorporate steps to solve the problem. For example, a patient may be able to calculate the number of nickels in \$1.00 and the number of nickels in \$0.35, but not the number of nickels in \$1.35.

Behavioral disturbance can occur with either cortical or subcortical dementia, but apathy and depression are more common in the early stages of a subcortical dementia and may predate other cognitive problems. The cortical signs, aphasia, apraxia and agnosia, are lacking in subcortical dementia and suggest either AD or a focal process such as stroke, mass lesion or focal degeneration (e.g. primary progressive aphasia).

Motor signs, such as increased tone and bradykinesia, are common in subcortical dementia. Abnormal movements, such as tremor, chorea or dystonia, may also occur. Posture is often abnormal and patients may appear stooped or extended when standing. Gait problems include poor initiation, small step length and difficulty turning. AD patients do not demonstrate motor signs, abnormal movements or posture and gait abnormalities until the latter stages of dementia.

DISEASES CAUSING SUBCORTICAL DEMENTIA

Extrapyramidal Disorders

Parkinson's Disease (PD)

PD is a common cause of subcortical dementia and is diagnosed by the clinical findings of bradykinesia, tone abnormalities (rigidity and cogwheeling) and rest tremor. Almost all patients with PD can be shown to display cognitive slowing and other features of subcortical dementia, but only a subset progress to frank dementia⁴. Further complicating matters, some demented patients with PD develop a mixed cortical and subcortical dementia. At autopsy, patients with mixed dementia may exhibit pathologic signs of AD and PD or diffuse, intraneuronal Lewy bodies. The exact relationship between Parkinson's dementia, Lewy body dementia and AD is poorly understood⁵.

Other parkinsonian syndromes are also associated with subcortical dementia. Progressive supranuclear palsy (PSP) can be distinguished from PD by limited voluntary vertical eye movements, absence of rest tremor, extended (rather than stooped) posture and lack of response to dopamine replacement⁶. Clinically significant dementia is more common in PSP than PD. Multiple systems atrophy (MSA) is another parkinsonian syndrome that can be associated with subcortical dementia. Patients with MSA have a variable combination of parkinsonism, cerebellar ataxia, corticospinal tract abnormalities and/or autonomic dysfunction⁷.

Huntington's Disease (HD)

HD is an autosomal dominant inherited neurodegenerative disease characterized by psychiatric abnormalities, chorea and subcortical dementia⁸. The earliest symptoms of HD are often psychiatric and demented HD patients display the classic features of subcortical dementia. HD is caused by pathologic expansion of a CAG repeat in the huntingtin gene. A blood test is commercially available to detect the causative mutation.

Cerebrovascular Disease

The location of infarcted brain tissue in a stroke determines its clinical features. Strokes involving the cerebral cortical gray matter will cause cortical signs such as aphasia and apraxia. Subcortical dementia occurs when infarction involves the deep gray matter nuclei (e.g. basal ganglia and thalamus) and/or the periventricular white matter⁹. For dementia to develop, damage must be bilateral and multifocal. Small vessel strokes can be caused by lacunar infarction or Binswanger's disease. Lacunar strokes occur when small intracerebral penetrating blood vessels are occluded and Binswanger's disease is the result of chronic white matter ischemia. Risk factors for small vessel cerebrovascular disease include cigarette smoking, hypertension and diabetes mellitus. Patients with small vessel cerebral infarction may not experience any traditional stroke-like episodes and deny a step-wise decline in cognitive function. The physical examination will, however, demonstrate focal neurologic signs, including pseudobulbar palsy, hyper-reflexia and pathologic reflexes, such as the Babinski sign. CT or MRI is useful in demonstrating small vessel strokes in patients with suspected subcortical dementia.

Miscellaneous Causes

Multiple sclerosis (MS) and HIV infection can cause subcortical dementia. These diseases are often suspected when confronted by a young person with dementia, but should also be considered in older patients. Dementia is usually a late finding in MS and is typically associated with other neurologic signs, including eye movement abnormalities, motor findings, sensory symptoms and/or cerebellar deficits. The presence of cerebellar ataxia and subcortical dementia is also seen in patients with spinocerebellar ataxias (SCA). There are a large number of different diseases that present as SCA and the diseases may be sporadic or inherited¹⁰. Magnetic resonance brain imaging of patients with advanced SCA will show cerebellar and/or brainstem atrophy. Genetic testing is available for many of the inherited forms of disease.

Hydrocephalus can cause subcortical dementia. Hydrocephalus may occur in the elderly without obvious lesions obstructing cerebrospinal fluid (CSF) flow and with normal CSF pressure [normal pressure hydrocephalus (NPH)]. Brain imaging of patients with NPH shows enlarged ventricles and widely patent intraventricular foramina. The clinical signs of NPH include the well-known triad of subcortical dementia, urinary incontinence and gait apraxia. CSF shunting of patients with NPH can result in significant improvement, but is typically unsuccessful if the dementia is long-standing or far-advanced. Other potentially treatable causes of dementia can also present, with a subcortical pattern including chronic infection and vasculitis.

Frontal lobe dementia presents as a classic subcortical dementia, despite the fact that the pathology is located in frontal cortical neurons. The subcortical-frontal neuronal pathways mediate many of the features of subcortical dementia, including apathy, inattention, bradykinesia and problems with planning^{3,11}. It is not surprising that damage to the subcortical nuclei, connecting white matter tracts or frontal neurons, present with a similar clinical phenotype. Other causes of frontal lobe damage can also cause a "subcortical" dementia, such as trauma, infarction, tumor or psychiatric disease.

CONCLUSION

Dividing the clinical presentation of patients with dementia into cortical and subcortical groups can be useful as a first attempt to identify the etiology of dementia, but this approach has limitations. Dementing diseases rarely cause purely cortical or subcortical pathology. Strokes can involve both white and gray matter. AD can cause degeneration in subcortical nuclei or frontal cortex and clinically resembles a subcortical dementia. Analogously, the pathology of frontal lobe dementia causes cortical neurons to degenerate, but the clinical profile is subcortical. Depression can further confuse the clinical picture by making a cortical dementia have subcortical features, leading to inaccurate diagnosis. The classical distinction between subcortical and cortical dementia is also less clear in patients with advanced dementia, because the clinical patterns tend to merge. Despite these problems, recognition of subcortical features can increase one's suspicion of unusual causes of dementia and lead to increased accuracy of diagnosis and, potentially, treatment.

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Early-onset Dementias

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The early-onset dementias are a heterogeneous group of neurodegenerative diseases with onset generally defined as prior to age 65. Research into these diseases has suffered from ascertainment bias, inconsistency in definitions, small numbers and inaccuracies associated with retrospective review of charts and death records. However, clinically useful information has gradually emerged over more than 60 years of study.

Liston reviewed the topic in 1979 and contributed a series of 50 cases^{1,2}. Despite several significant advances in the field since that time, most notably the discovery of the presenilin genes, his review continues to be an excellent starting point in understanding this group of diseases.

EPIDEMIOLOGY

Early-onset dementia represents approximately 5–10% of all dementias seen by health caregivers.

The relative frequency of common dementia subtypes in early-onset dementia roughly parallels that for senile dementia. The most prevalent form of dementia is Alzheimer's disease (AD), followed by vascular dementia (VaD). In countries with a high rate of alcohol use, alcoholic dementia has a frequency roughly equal to vascular dementia. The frequency of less common dementing illnesses, such as inborn errors of metabolism and chronic CNS infection, is expected to be higher than in older age groups; however, epidemiological studies are lacking.

Estimates of annual incidence for early-onset AD (EOAD) vary from 2.4 to 22.6/100 000^{3–7}. One study found a point prevalence of 34.6/100 000 with a 5 year survival of 68%. Median survival was estimated as 8.1 years in another study. These estimates suggest a more aggressive course for EOAD compared with the senile variant. Most studies support increased risk among females for PSAD, with a relative risk of 1.0–1.7.

Only a few epidemiological studies have attempted to include other etiologies of dementia. Two studies from Scotland varied widely with respect to relative frequencies of PSAD and vascular dementia^{3,6}. One study found comparable frequencies, while another found the frequency of vascular dementia to be about 20% that of PSAD. In the latter study, of 114 patients, 60 had PSAD, 13 had vascular dementia, 14 had alcohol-related dementia, 25 had overlapping diagnoses and two had other diseases.

EARLY-ONSET ALZHEIMER'S DISEASE

EOAD differs from senile onset AD in several respects, including patterns of inheritance and disease course⁸. While

Table 50c.1 Genes linked to familial EOAD

Gene	Chromosome	Known mutation as of 1999
Amyloid precursor protein (APP)	21	5
Presenilin 1	14	45
Presenilin 2	1	2

the majority of cases of EOAD are sporadic, there is a significantly increased frequency of familial clusters in comparison to senile AD⁹. These clusters have facilitated linkage analysis, leading to the identification of a number of mutations in several genes. So far, three genes have been linked to familial EOAD (Table 50c.1). It is estimated that approximately 50% of familial cases of EOAD can be attributed to known mutations, suggesting that other genes yet to be identified are involved as well.

The mutations listed in Table 50c.1 all result in increased abnormal cleavage of APP, ultimately leading to production of amyloid plaques. The inheritance pattern for mutations in all three genes is autosomal dominant with high penetrance. No practical screening tests are currently available.

The apolipoprotein E (ApoE) gene on chromosome 19 has been identified as an important susceptibility gene for EOAD. Three naturally occurring allotypes of ApoE are known, ApoE 2, 3 and 4. Individuals homozygous for ApoE 4 have approximately twice the relative risk of developing AD as individuals homozygous for ApoE 3. Conversely, individuals with one or two copies of ApoE 2 have approximately half the risk for EOAD as individuals homozygous for ApoE 3. Risk appears to be greatest in individuals developing cognitive impairment between 60 and 65 years of age and declines in older age groups.

The role of ApoE in AD is poorly understood and is the topic of intensive research. In addition to its role in AD, homozygosity for ApoE 4 has been found to be a risk factor for poor outcome following stroke. Thus, ApoE is thought to possibly play a role in mediating inflammatory responses in the brain.

Age at onset of AD has been shown to be an important determinant of disease course and clinical presentation, with more rapid cognitive and functional decline observed in EOAD. On neuropsychological testing, individuals with EOAD score significantly worse on attentional items compared with senile AD. These observations have suggested the possibility that EOAD may be a different clinical entity than senile AD.

VASCULAR DEMENTIA

Little information exists regarding early-onset vascular dementia (EOVD), although the relative frequency in some areas, notably Japan, appears to be equal to EOAD. Risk factors are similar to senile vascular dementia and include hypertension, diabetes mellitus, smoking and hypercholesterolemia. As with EOAD, there is an increased prevalence of familial clusters in EOVD, most of which can be attributed to homocysteinuria, although other inherited coagulopathies may play a significant role. There is probably a considerable overlap between cases of EOAD and EOVD. Readers interested in additional information about the diagnosis of AD or VaD are referred to reviews by Gersing *et al.*¹² and by Doraiswamy *et al.*¹³

APPROACH TO DIAGNOSIS

There are perhaps several hundred disorders with dementia as a clinical feature. Most of these disorders are quite rare. This diversity in etiology poses a significant challenge to the clinician. For example, a case report of two patients with clinically similar features found one to have EOAD and the other to have adult onset metachromatic leukodystrophy¹⁰.

The following general guidelines, although not exhaustive, are intended as an aid in the work-up of patients presenting with cognitive impairment at age 65 and younger:

1. History of present illness:
 - (a) A gradual insidious course is suggestive of EOAD.
 - (b) Cognitive decline in association with a stroke or "spells" raises the index of suspicion for EOVD.
 - (c) A lifelong history of underachievement, childhood delay in milestones or difficult pregnancy or delivery is suspicious for an inherited disorder affecting cognition.
2. Ask about risk factors for less common dementias:
 - (a) Exposure to toxins, including carbon monoxide, solvents, pesticides, heavy metals.
 - (b) History of childhood encephalitis or meningitis.
 - (c) History of multiple concussions.
 - (d) History of malabsorption, diarrhea or significant change in diet.
 - (e) Parental age at birth.
 - (f) Detailed history of alcohol and drug use.
 - (g) AIDS risk factors.
3. Past medical history:
 - (a) Search for potentially reversible causes of dementia:
 - (i) Thyroid disease.
 - (ii) Epilepsy.
 - (iii) Syphilis.
 - (iv) Pernicious anemia.
 - (v) Lupus, sarcoidosis, other CNS inflammatory diseases.
 - (vi) Cancer.
 - (vii) Hepatitis.
4. Search for illnesses that may mimic dementia:

Table 50c.2 Checklist for neurological examination in dementia diagnosis

Dementia associated with neuropathy	3. Alcoholic dementia
1. Hypothyroidism	4. Paraneoplastic encephalopathy
2. Nutritional disorders, especially B ₁₂ deficiency	5. Hydrocephalus
3. Vasculitis, especially Sjogren's syndrome, SLE	Dementia associated with dysautonomia
4. Exposure to solvents and alcohol	1. Multisystem atrophy
5. Multi-infarct dementia associated with diabetes	2. Multi-infarct dementia with diabetes
6. HIV	3. Parkinson's disease
7. Neurosyphilis	Dementia associated with myoclonus
8. Lyme disease	1. Early-onset fronto-temporal dementia
9. Metachromatic leukodystrophy	2. Creutzfeldt-Jakob disease
10. Adrenoleukodystrophy	3. Post-anoxic encephalopathy
Dementia associated with hemiparesis	4. Autoimmune thyroiditis
1. Multi-infarct dementia	5. Whipple's disease
2. Mitochondrial encephalomyelopathy with stroke-like episodes (MELAS)	Dementia associated with epilepsy
3. Vasculitis from various causes	1. Metastatic brain lesions
4. Masses in the brain, including tumors, vascular malformations, hematomas, abscesses, etc.	2. Cryptococcal meningitis
5. Complex partial epilepsy	3. Dentato-pallido-luysian atrophy
6. Amyotrophic lateral sclerosis	4. Anti-epileptic drug-related encephalopathy
Dementia associated with tremor or rigidity	Dementia associated with eye movement abnormalities
1. Early-onset Parkinson's disease	1. Multiple sclerosis (intranuclear ophthalmoplegia)
2. Multisystem atrophy	2. Progressive supranuclear palsy (impaired vertical gaze)
3. Spino-cerebellar degeneration	Dementia associated with myopathy
4. Multi-infarct dementia, especially with involvement of the basal ganglia	1. Myotonic dystrophy
5. Huntington's disease	2. Hypothyroidism
6. Wilson's disease	3. Diabetes mellitus
7. Progressive supranuclear palsy	Dementia with variable physical findings
8. Hallevorden-Spatz disease	1. Primary HIV dementia
9. Manganese poisoning	2. PML
10. Post-encephalitic dementia	3. ADEM
11. Multiple sclerosis	4. Metabolic causes, e.g. hypothyroidism, hypercalcemia, Addison's disease
12. Carbon monoxide poisoning	5. Dementia with Lewy bodies
13. Cryptococcal meningitis	Dementia with a positive family history
14. Dementia pugilistica	1. Familial AD (Presenilin 1 and 2 mutations)
Dementia associated with nystagmus and/or ataxia	2. Inborn errors of metabolism
1. Wernicke's encephalopathy	3. Huntington's disease
2. Spino-cerebellar degeneration	4. Myotonic dystrophy

- (a) Sleep disorders, including sleep apnea and narcolepsy.
 - (b) Depression, PTSD, anxiety disorders.
 - (c) Ongoing occult substance abuse.
 - (d) Subacute delirium.
5. Review medications:
- (a) Long-term anticholinergic, neuroleptic and/or anti-convulsant use is associated with cognitive impairment.
 - (b) Overaggressive treatment of hypertension can result in hypoxic/ischemic encephalopathy.
6. Family history:
- (a) Strokes or MI at an early age suggests coagulopathy or familial hypercholesterolemia.
 - (b) Strong family history of diabetes mellitus should prompt search for occult diabetes.
 - (c) Early-onset dementia without other neurological findings suggests familial EOAD.
 - (d) Amyotrophic lateral sclerosis has a familial association with frontotemporal dementia.
7. Mental status examination is intended to reveal the pattern of cognitive impairment and should minimally include:
- (a) Mini-Mental Status Examination.
 - (b) Additional tests for retrieval, such as list generation.
 - (c) Additional tests for attention, such as asking to recite months in reverse.
 - (d) Additional tests for visuospatial function, such as drawing a transparent cube.
 - (e) Tests of motor praxis, such as demonstrating untying and tying a shoelace.
 - (f) Tests of frontal lobe function, such as Luria sequencing, visual go-no go testing and general assessment of judgment.
8. Thorough neurological examination:
- (a) See Table 50c.2.
9. Head imaging:
- (a) Some form of head imaging should probably be performed in all cases of early-onset dementia, especially if focal neurological findings are present.
 - (i) While CT is readily available and inexpensive, MRI is superior in imaging the posterior fossa and in distinguishing between demyelination and edema in white matter lesions.
10. Laboratory studies:
- (a) Should be tailored to the findings on history, mental status and physical examinations.
 - (i) The interested reader is referred to an excellent and exhaustive review of rare dementia syndromes by Reichmann and Cummings¹¹.
 - (b) All patients should be tested for blood count, electrolytes, liver panel, albumin, B₁₂, folate, VDRL, sedimentation rate.
 - (i) While the yield of these investigations is low, their cost is low as well, and they screen for reversible causes of dementia.
 - (c) ApoE genotype is reasonable in cases suspicious for EOAD, especially if there is no significant family history.

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Creutzfeldt–Jakob Disease and Other Degenerative Causes of Dementia

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The following disorders may have dementia as a prominent part of the illness, but overall they are not common in clinical practice. Rarely is it possible to differentiate them on the basis of the form that the dementia takes, but other features can make them sufficiently distinctive to be worthy of description.

CREUTZFELDT–JAKOB DISEASE (CJD)

This disorder was first described in 1921^{1,2}, when an unusual fatal neurological illness in six cases was associated with a microscopic spongy appearance of the brain at post mortem. It is a rare, rapidly progressive dementing illness, affecting both sexes, with an onset usually after the age of 55, the exceptions being those cases associated with transmission from human-derived products or tissues, or new variant CJD. The presenting features include myoclonus, cortical blindness, pyramidal/extrapyramidal/cerebellar features and akinetic mutism. It is universally fatal, with an illness duration that rarely exceeds 6 months. No cause has been found, but the existence of similar neurological illnesses in other mammals also with spongy appearances of the brain has led to much research.

In the 1950s, a disease known as kuru was discovered in a tribe that practised cannibalism in the New Guinea highlands. This had many clinical and pathological similarities to CJD, and in 1968 the disease was transmitted to monkeys by intracerebral inoculation of extracts of kuru brain³. Kuru was felt to have been transmitted to other members of the tribe through their practice of eating part of the remains of a previous kuru victim, especially the brain.

Scrapie is a spongiform encephalopathy of sheep that has been recognized for over 250 years and has been known to be transmissible to other sheep by natural means, both vertically and horizontally, since the 1930s. In 1986, cattle were also found to have developed a bovine spongiform encephalopathy (BSE)⁴. The theory advanced to account for this was that they had been exposed to scrapie through the consumption of rendered sheep remains as part of their diet.

An altered form of a normal brain protein has been found in cases of spongiform encephalopathy in both man and animals, which shows resistance to protease digestion and is known as protease-resistant protein (PrP) or “prion” protein⁵. The normal variant is encoded for by a gene on chromosome 20 in humans⁶. Already a number of mutations of this gene in cases of CJD have been discovered⁷, and familial cases are invariably associated with abnormalities of the gene. Homozygosity for

methionine at the site of a common polymorphism at codon 129 on the PrP gene is found with twice the normal incidence in cases of sporadic CJD⁸, whereas valine homozygosity incidence is higher in cases associated with human growth hormone administration.

The possibilities of the disease being due to a viral or immunological process has largely been discounted, and current opinion is focusing on somatic mutation in the PrP gene as a possible cause.

The incidence of CJD, with few exceptions, is remarkably constant worldwide, at approximately 0.5–1/million population/year. Clustering of cases has been recognized⁹ and this is thought to represent genetic susceptibility, rather than the presence of an environmental factor. Sporadic CJD is not felt to be due to transmission from consumption of or contact with animal products¹⁰. Cases have occurred in recipients of human-derived products such as growth hormone, gonadotrophin, corneal grafts and dura mater grafts, and following neurosurgical procedures^{11–14}. Notable amongst these have been cases in patients who received human pituitary extracts, in whom pathologically confirmed CJD was manifest 10–20 years later, suggesting a prolonged “incubation period” before the disease occurs.

Diagnosis of CJD can only be made with certainty at post mortem or by brain biopsy. However, the diagnosis has been confirmed by tonsillar biopsy in suspected cases¹⁵, although this technique is not widely recommended in view of the possible risk to operators¹⁶. The finding of elevated levels of cerebrospinal fluid levels of S100 and 14-3-3 proteins is relatively specific for suspected CJD, although raised levels are also found in encephalitis¹⁷.

The rapidity with which the dementia develops, with global cortical and subcortical components as well as the specific signs mentioned earlier, should avoid confusion with other causes of dementia. The electroencephalogram (EEG) often shows a typical appearance of generalized repetitive triphasic complexes, with virtual abolition of background rhythms, but this feature is not invariable, often occurring late in the course of the illness. Its absence in no way excludes the disorder. There is almost always a slowing of cortical rhythms into the delta range. Computed tomography (CT) scan is usually normal, but is useful to exclude other conditions. Magnetic resonance imaging (MRI) can show non-specific high signal on T2-weighted images in the basal ganglia in some cases.

There is no treatment available, although there have been isolated case reports of slowing of the rate of progression of the disorder with amantadine¹⁸.

NEW VARIANT CJD

The first cases of new variant CJD appeared in 1995, when individuals under the age of 40 years developed a rapidly progressive dementing illness, and pathological analysis at post mortem confirmed the typical features of spongiform encephalopathy¹⁹. There were a number of important differences to typical sporadic CJD, however:

1. Many of the cases presented with features of a psychiatric disorder, such as anxiety, depression or hysteria.
2. The development of the dementia was more slowly progressive than in typical cases of CJD.
3. The typical changes of triphasic complexes on the EEG were lacking.
4. Brain microscopy revealed far more extensive PrP-positive plaques.

The evidence that these cases may be directly linked to BSE in cattle is compelling in that the PrP protein in these cases is biochemically very similar to that found in cases of BSE, and the incubation time of transmitted cases in transgenic mice expressing the human PrP gene is identical to that of transmitted BSE cases²⁰. Up to September 2001, 107 cases had been described, one of which was in an elderly man aged 74.

MULTISYSTEM ATROPHY (SHY-DRAGER SYNDROME), PROGRESSIVE SUPRANUCLEAR PALSY (PSP) (STEELE-RICHARDSON-OLSZEWSKI SYNDROME), DENTATOPALLIDO-LUYSIAN ATROPHY AND OLIVOPONTocerebellar ATROPHY (OPCA)

Dementia is not an invariable accompaniment of any of the above, but thorough cognitive testing will often reveal deficits, especially in late cases. The dementia may be of frontal lobe type or sub-cortical but not severe, and may be masked by other clinical features that are more prominent²¹.

Nomenclature of the above conditions is subject to debate. Extrapyrarnidal features are seen in most, although a clinical presentation with prominent cerebellar features tends to result in a patient being labelled as OPCA and pronounced autonomic failure as Shy-Drager syndrome, but the pathological changes of cell loss and gliosis in all are similar and may also be present in the same sites, although to varying degrees. Only in PSP is there a clearly distinct clinical presentation and histological appearance.

The early signs in all the above conditions are usually rigidity in muscle tone, bradykinesia of movement and postural instability. The signs are usually bilateral and progress over months and years.

In PSP, first described in 1964²², there is characteristically a loss of conjugate voluntary eye movements, beginning with vertical gaze. The range of eye movement is improved if the patient is made to fixate on a target and the head moved. Upper motor neurone limb signs are common, and dystonic posturing of neck muscles resulting in extension is often seen. Reduction in verbal fluency is a prominent feature of the condition, and it has been postulated that this is due to interruption of fronto-basal circuitry²³. The condition is confirmed pathologically by the finding of neuronal loss, neurofibrillary tangles and gliosis primarily affecting the subthalamic nucleus, globus pallidus, dentate, substantia nigra, locus coeruleus, periaqueductal grey matter and other brainstem nuclei.

Multisystem atrophy (MSA) results in a degenerative process affecting neurones throughout the central nervous system (CNS) and thus the signs may be widespread, with involvement of corticospinal tracts and especially the autonomic system. Diag-

nosis is usually clinical and may be strongly suspected when there are additional signs, such as laryngeal stridor or denervation of the urethral sphincter on electromyography.

Dentatorubral-pallido-luysian atrophy is an autosomal dominant disorder that has been found to be linked with a trinucleotide repeated on the B37 gene of chromosome 12. The clinical features include seizures, chorea, dementia, ataxia, mental retardation and psychiatric disease. Abnormalities are seen in the subcortical white matter.

Other investigations in this group of conditions are usually normal, although evidence of cortical atrophy may be present in late cases on the CT scan, and a case has been made for the differentiation of PSP from other causes of extrapyramidal syndromes and dementia by subtle findings on CT²⁴.

The parkinsonian features in these disorders are often resistant to treatment with conventional anti-parkinsonian drugs. In the case of MSA, L-dopa therapy can exacerbate symptoms of postural hypotension due to the coexistent autonomic neuropathy. The disorders are therefore treated along supportive lines, although MSA may require more specific drug therapy, directed at features such as postural hypotension.

In all the above disorders, depression can be a common accompanying symptom, as patients are aware of the restriction in activity caused by the disease. This may require separate treatment.

As mentioned earlier, a dementing illness may also occur, especially in the later stages. However, this may be overlooked if poor performance on a task is attributed to slowness of response or the effects of drug therapy.

MOTOR NEURONE DISEASE

This disorder, which primarily causes loss of both upper and lower motor neurones, can also be associated with a dementia. Dementia is detectable in approximately 5% of cases, although its existence may be obscured if the patient is rendered anarthric and paralysed. It may precede the onset of the typical signs of motor neurone disease in about 50% of cases, and is more common in those with a bulbar onset or in familial cases. Myoclonus has been noted in up to 15% of cases and this is probably what has been responsible for the confusion with Creutzfeldt-Jakob disease—with many of these cases being previously labelled as the “amyotrophic form of CJD”, although the other clinical features of CJD, the typical EEG findings and laboratory transmission to animals, were lacking²⁵. The dementia is typically “frontal” in type, with deficits in attention, learning, naming, insight and judgement²⁶.

ALS may also occur in association with both parkinsonism and dementia, with the dementia being indistinguishable from that occurring in Alzheimer's disease. The age of onset was almost the same for the parkinsonism, ALS and dementia, indicating that they are probably of the same aetiology.

The studies in Guam²⁷ have shown the development of a parkinsonism-dementia-ALS disease on exposure of individuals to the toxins from the cycad plant. In the review by Hudson²⁸, it is proposed that this link between the three modes of presentation may indicate that the disorders may be variants of the same disease and not unique to Guam.

NORMAL PRESSURE HYDROCEPHALUS

The title of this condition is a misnomer, as the cerebrospinal fluid (CSF) pressure is raised, but only intermittently, to produce what are known as “B” waves on continuous CSF pressure recordings. However, classic symptoms of raised intracranial pressure are

absent, the patient presenting with a triad of symptoms comprising dementia, gait dyspraxia and urinary incontinence. Cerebral atrophy is seen on CT scan, but the condition may be suspected when the ventricles are disproportionately large in comparison with the degree of sulcal widening. Unfortunately there is no reliable diagnostic test, as evidenced by recent reports in the literature²⁹. Some claims have been made for isotope cisternography, which shows delayed passage of radioactivity from the ventricles to the area surrounding the cerebral convexities. Treatment is by the insertion of a CSF shunt, and in some cases this has even been used as the definitive diagnostic test. No cause has been found to account for the condition, although in some cases a "secondary" form is recognized as being due to previous meningeal inflammation or head injury and subsequent impairment of CSF uptake.

POST-TRAUMATIC DEMENTIA

Dementia pugilistica in boxers presents as a progressive neurological disease with involvement of pyramidal, extrapyramidal and cerebellar systems, in addition to memory loss and personality change. The onset may be many years after the cessation of a boxing career. The pathological appearances resemble those of Alzheimer's disease, but neurofibrillary tangles predominate, with an absence of plaques³⁰.

HUNTINGTON'S DISEASE

This autosomal dominant inherited condition, with an onset usually after the age of 30 and rarely after the age of 70, has choreiform movements and dementia as its most common presenting features. Diagnosis can now be confirmed by analysing the huntingtin gene on chromosome 5 for CAG triplet repeats in excess of 35. Atrophy of the caudate nucleus may be seen in typical cases on CT or MRI scan. The dementia does differ from that of Alzheimer's disease by virtue of the fact that there is more difficulty with tests of letter fluency and the copying of geometric features³¹, indicating that the dementia is of the subcortical type.

FAMILIAL SPASTIC PARAPARESIS, OR SPINOCEREBELLAR ATROPHY ASSOCIATED WITH DEMENTIA

These conditions are considered together as both may produce a dementing-type illness which is in itself indistinguishable from other forms of dementia, but is set apart by the presence of other signs, such as a spastic paraparesis or a cerebellar syndrome. In both, the dementia advances only slowly, and may only be a minor feature of the illness, although recent work has shown that the cognitive impairment may be subclinical in some kindreds and overlooked with routine clinical screening tests³².

PROGRESSIVE SUBCORTICAL GLIOSIS

This condition, described in 1967, presents as a dementing process with subcortical features and may mimic Alzheimer's disease. Infrequent findings are signs of extrapyramidal involvement, and cases with supranuclear gaze palsy have been described³³. The clinical course extends over years, often with a final akinetic mute stage. The cardinal pathological feature is intense gliosis and astrocytic hyperplasia affecting the subcortical structures of the thalamus, basal ganglia, brainstem grey matter and ventral horns of the spinal cord.

CORTICO-BASAL DEGENERATION

The more prominent signs of extrapyramidal dystonias, cortical sensory loss, dyspraxia, myoclonus (often focal) and the "alien hand" sign, tend to obscure a slowly developing dementia which is usually cortical in nature. The presence of ideomotor dyspraxia may help to distinguish this extrapyramidal syndrome from disorders such as PSP³⁴. The mean age of onset in a series of 15 patients was 60 years³⁵. No investigation is diagnostic, and confirmation of the diagnosis is by finding nerve cell loss and gliosis in the frontoparietal cortex and extrapyramidal system.

DEMENTIA LACKING SPECIFIC HISTOLOGICAL FEATURES

As well as being without specific histology, the pattern of dementia is unremarkable, being of a frontal lobe type more closely resembling Pick's disease, with many cases also showing extrapyramidal features. The pathological features are cortical vacuolation and astrocytosis in the deeper layers, without neurofibrillary tangles, plaques or Pick or Lewy bodies.

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Frontotemporal Dementia (Pick's Disease)

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“Frontotemporal dementia” is the term currently preferred to describe patients with focal cortical atrophy involving the frontal and/or temporal lobes. Pathologically, such patients show severe neuronal loss, spongiosis and gliosis, and in a minority of cases classic argyrophilic, tau-positive intraneuronal inclusions are present (Pick bodies). The hallmark changes of Alzheimer's disease are virtually always absent¹.

One of the most exciting developments in the past decade has been the discovery of mutations in the tau gene on Chromosome 17 in some families with dominantly inherited FTD², although it should be stressed that most cases are sporadic. Clinically, FTD patients present with one of three major syndromes, which reflect the initial locus of pathology: dementia of frontal type, semantic dementia and progressive non-fluent aphasia³. It should be noted that each of these syndromes can be associated with motor neurone disease and a full neurological evaluation should also be included, especially in rapidly progressive cases and if bulbar symptoms develop⁴.

Patients with FTD present below the age of 65 and there is an equal sex distribution. Although rare, FTD is the second commonest cause of dementia in the presenium (after Alzheimer's disease)⁵. In Cambridge we have studied approximately 100 cases with FTD; the rarest syndrome is non-fluent progressive aphasia, the other two account for approximately 40% of cases each. The average survival from diagnosis is, on average, 10 years. In our experience, patients with the frontal form of the disease progress at the slowest rate, semantic dementia cases have a more rapid course and those with motor neurone disease-associated FTD rarely survive more than 2 years.

DEMENTIA OF FRONTAL TYPE (DFT) (FRONTAL VARIANT FTD)

The onset of symptoms is insidious and insight is lacking. Relatives complain of a change in personality and behaviour: disinhibition, poor impulse control, antisocial behaviour, stereotypical features (e.g. insisting on eating the same food at exactly the same time daily, cleaning the house in precisely the same order or the use of a repetitive catchphrase) and a change in appetite and food preference towards sweet food are the features that best discriminate FTD from Alzheimer's disease. They reflect the early involvement of the orbitobasal frontal cortex^{6,7}. Apathy is also very common but non-specific. Deficits in planning, organization and other aspects of executive function are universal as the disease progresses to involve the dorsolateral prefrontal cortex. Major depression and psychosis are rare.

A major advance in the area has been the development of a semi-structured carer interview, the Neuropsychiatric Inventory, which appears to be able to differentiate patients with FTD and AD⁸.

Neuropsychological Findings

Some patients show clear-cut cognitive deficits at presentation but most traditional “frontal executive” tasks are sensitive to dorso-lateral, rather than orbitobasal frontal, dysfunction; among the most useful are the Wisconsin Card Sorting Test and verbal fluency (i.e. the generation of words beginning with a given letter of the alphabet). Recently, quantifiable tasks involving decision-making and risk-taking and better able to detect orbitobasal frontal function have been developed⁹.

Memory is relatively spared: orientation and recall of personal events is good but performance of anterograde memory tests is more variable, and patients tend to do poorly on recall (as opposed to recognition)-based tasks. A reduction in spontaneous conversation is common, but patients with DFT perform well on tests involving picture naming and other semantically-based tasks. Visuo-spatial abilities are strikingly preserved, particularly when the organization aspects are minimized: the Rey figure test is often copied poorly due to impulsiveness and poor strategy¹⁰.

Simple cognitive screening tests, such as the Mini-Mental State Examination (MMSE), are unreliable for the detection and monitoring of patients with DFT, who frequently perform normally even when requiring nursing home care¹¹.

SEMANTIC DEMENTIA (TEMPORAL LOBE VARIANT FTD)

Patients with this variant of FTD present with a progressive fluent aphasia but the underlying cognitive deficit is a breakdown in semantic memory^{3,12}. Semantic memory is the term applied to the component of long-term memory which contains the permanent representation of things in the world, including objects, words and people. It is the database which gives meaning to our sensory experiences.

Patients complain of “loss of memory for words”. Although aware of their shrinking expressive vocabulary, patients are strangely oblivious to their impaired comprehension. Since the grammatical and phonological structure of language remains intact, the changes are relatively subtle, at least in the early stages. Patients with predominant right-sided atrophy may present with difficulty recognizing faces (prosopagnosia), at first affecting less commonly encountered people but with time severe prosopagnosia

develops: in contrast to the true modality-specific prosopagnosia, which occasionally complicates right occipitotemporal strokes, the deficit in semantic dementia is cross-modal, so that patients are also impaired in the identification of names and voices¹³.

In contrast to Alzheimer's disease, patients with semantic dementia are well orientated and have good episodic (day-to-day and autobiographical) memory, although recent studies have shown that the preservation applies only to recent memories. That is to say, they show a reversal of the temporal gradient found in amnesia and Alzheimer's disease¹⁴.

Behavioural changes may be slight at presentation but with time features identical to those seen in DFT emerge and may be striking in patients with right-sided disease¹⁵.

Neuropsychological Findings

The impairment of knowledge is most apparent on tasks requiring a verbal output, such as category fluency tests (in which subjects are asked to produce as many examples as possible from defined semantic categories, such as animals, within 1 minute), picture naming and the generation of verbal definitions to words and pictures. The pattern of errors reflects a loss of fine-grained knowledge, with preservation of superordinate information: on naming tasks, naming errors are initially category co-ordinates ("elephant" for hippopotamus), then with time prototype responses emerge, so that all animals are called "dog", then eventually they are simply called "animal". Single-word comprehension is also affected, as judged by tasks such as word-picture matching or synonym tasks (e.g. which of the following is the odd one out, "pond, lake or river"). Non-verbal semantic knowledge is less easy to assess, but the Pyramids and Palm Trees Test in which the subject is asked to judge the semantic relatedness of pictures, invariably reveals deficits^{10,16}.

In contrast to the profound semantic deficit, other aspects of language competency (phonology and syntax) are strikingly preserved. Although able to read and spell words with regular spelling-to-sound correspondence, virtually all cases have difficulty in reading and spelling irregular words (e.g. reading PINT to rhyme with hint, flint, etc.). This pattern, known as surface dyslexia (or dysgraphia), has been attributed to the loss of semantic support which is necessary for the correct pronunciation of irregular words. Patients perform normally on non-verbal problem-solving tasks, such as Raven's Matrices, and on tests of perceptual and spatial ability.

NON-FLUENT PROGRESSIVE APHASIA

The status of patients with the non-fluent form of progressive aphasia within the spectrum of FTD is less certain. Changes in behaviour are rare, but after a number of years global cognitive decline occurs. Unlike the other syndromes, a number of non-fluent cases have Alzheimer pathology¹⁷.

Patients present with complaints of speech dysfluency and distortion or word-finding difficulty. Phonological errors are usually obvious in conversation. Comprehension is relatively well preserved, at least in the early stages, although as the disease progresses there are problems with phoneme discrimination, which the patients invariably attribute to poor hearing. In the late stages patients become mute and effectively "word deaf". Day-to-day memory is good and patients cope well in everyday life.

Neuropsychological Findings

The pattern of cognitive deficits in progressive non-fluent aphasia is the mirror image to that found in semantic dementia. They

perform well on tests of semantic memory, except on those requiring a spoken output. Although conversational speech is severely disrupted, the anomia is mild and the errors are phonological (elegant for elephant). Semantic category fluency is less affected than letter fluency. Word-picture matching, synonym tasks and other semantic tests are usually performed perfectly. They perform poorly on tests of phonological competence (such as repetition of multisyllabic words and rhyming) and syntactic comprehension¹⁶. In common with the other syndromes, however, performance on visuo-spatial and perceptual function is well preserved.

NEURORADIOLOGICAL FINDINGS IN FTD

CT scans are of limited value in diagnosis: the temporal lobes are poorly seen due to bone artefacts but MRI with coronal images is extremely valuable, especially in cases with semantic dementia who show focal atrophy of the polar region, fusiform and infero-lateral gyri with relative sparing of the hippocampal formation. An asymmetric pattern is almost invariable, with the left temporal lobe much more often involved than the right¹⁸. In non-fluent cases the changes are more subtle, with widening of the left Sylvian fissure. In dementia of frontal type, atrophy of the orbitofrontal cortex can be seen, but ⁹⁹Tc-HMPAO-SPECT is more valuable: focal hypoperfusion is apparent before clear structural changes are obvious^{11,19}.

MANAGEMENT

The management of patients and their families requires a specialist multidisciplinary team approach with input from clinical psychology, genetics, psychiatry and neurology. Counselling, especially with regard to genetic implications and prognosis, are essential at an early stage. Although there are, at present, no therapies that will affect the course of the disease, many of the symptoms can be helped. The stereotypical features, disinhibition and overeating, may respond to serotonin reuptake blockers²⁰. Neuroleptic drugs may be necessary to control aggressive behaviour.

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Alcoholic and Other Toxic Dementias

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ALCOHOLIC DEMENTIA

The concept of a dementia consequent upon the effects of long-term alcohol abuse developed from clinical observations of a gradual deterioration in personality and intellect in many alcoholics. Recent studies suggest that this is age-related, is milder in degree than the neurodegenerative dementias and can be present in 11–24% of demented patients^{1–3}. Alcoholic dementia was originally considered to be distinct from the amnesia of the Wernicke–Korsakoff syndrome, which also occurs in alcoholics but is caused by thiamine malnutrition rather than alcohol neurotoxicity. Horvath⁴ prospectively examined 100 chronic alcoholics presenting with a dementia syndrome and concluded that “alcoholic dementia” is not synonymous with the Wernicke–Korsakoff syndrome, and that other non-amnesic organic syndromes exist in alcoholics, characteristic of frontal, parietal or global cortical damage. Cutting⁵ performed a retrospective case-note analysis of alcoholic patients with cognitive impairment and found two forms of clinical presentation. One consisted of a rapidly developing illness in younger patients with preserved intellect, akin to the traditional Wernicke–Korsakoff syndrome, whereas the other was more characteristic of dementia, being a gradual and global cognitive deterioration in older patients.

This concept was soon challenged by the results of clinicopathological studies. Torvick *et al.*⁶ examined the clinical records of patients who, at autopsy, had diencephalic lesions characteristic of thiamine malnutrition. Of these, 75% were considered to be demented in life rather than amnesic and the majority had no additional neuropathological hallmarks of a neurodegenerative dementia. Torvick *et al.* concluded that the diencephalic lesions of thiamine deficiency can result in the clinical pictures of both “alcoholic dementia” and the Wernicke–Korsakoff syndrome. Victor and Adams⁷ then pointed out that 10% of their pathologically-proven cases of the Wernicke–Korsakoff syndrome developed cognitive abnormalities insidiously, rather than acutely⁸, and that cognitive impairments other than amnesia, and behavioural abnormalities such as inertia and apathy, can be demonstrated in these patients⁹. Thus, they argued that, depending on the severity of the non-mnemonic deficits or the mode of presentation, cases of the Wernicke–Korsakoff syndrome may be misattributed as cases of alcoholic dementia. The neuropathological studies of Harper and colleagues^{10,11,62} are in agreement with this. They found that two-thirds of the alcoholics coming to post mortem in their unit had lesions of thiamine deficiency, yet only one-third of these had received a clinical diagnosis of Wernicke–Korsakoff syndrome in life. In the remainder, the most common diagnosis was dementia.

Thus far, the evidence points to the conclusion that subcortical lesions caused by thiamine malnutrition can be sufficient to

explain both amnesia and dementia witnessed in alcoholics. However, Victor and Adams⁷ and Torvick *et al.*⁶ did not assume that *all* cases of alcoholic dementia are unrecognized cases of the Wernicke–Korsakoff syndrome. Both considered that superadded cerebral lesions may explain the dementia-like presentation of a proportion of patients with the Wernicke–Korsakoff syndrome. Because these additional lesions can be attributed to a variety of pathological processes, including chronic hepatocerebral degeneration, communicating hydrocephalus, Alzheimer’s disease and ischaemic infarction, they argued that there is no need to invoke a special process of alcohol neurotoxicity. More recent research supports this contention. Kasahara *et al.*² compared young (35–45 years) and old (>60 years) alcoholics and found evidence of dementia only in the older group. Most cases had additional medical diagnoses, including hypertension, liver disease and cardiomyopathy, and no case of dementia could be accounted for by the direct effect of alcohol.

Although dementia in alcoholics is unlikely to be caused solely by alcohol neurotoxicity, there is compelling evidence to suggest that alcohol is neurotoxic. Carefully controlled neuropathological studies have frequently found whole brain atrophy, predominantly involving the white matter, in chronic alcoholics^{12–16}. Neuronal death has been found in specific areas of the frontal association cortex^{17–19} and neuronal shrinkage in the cingulate and motor cortex^{17,20,21}. The contribution of liver failure to this neuropathology has been assessed in several studies and the bulk of evidence suggests that cirrhosis alone does not account for the brain shrinkage witnessed in alcoholics^{13,15,22}. However, existing studies of the contribution of thiamine malnutrition to such findings are equivocal with evidence that thiamine deficiency both does and does not cause the cortical damage seen in alcoholics^{13,15}. Medial temporal lobe limbic structures, i.e. the hippocampus and the amygdala, have been foci of interest because of their involvement in cognitive function. Reduced volumes have been reported in alcoholics but, again, it is not clear whether it is alcohol *per se* or thiamine deficiency that accounts for this^{16,19,23–25}.

It can be concluded from these studies that: (a) alcohol itself can cause neuronal damage; (b) liver disease *per se* is not a major factor in the aetiology of the neuropathological changes; and (c) thiamine malnutrition potentiates the neurotoxic effect of alcohol on the brain. Butterworth²⁶ proposes mechanisms to explain these conclusions. He argues that thiamine deficiency is common in alcoholism because of poor diet and gastrointestinal disorder. Alcohol and its metabolite acetaldehyde are directly neurotoxic and have toxic effects on thiamine-dependent enzymes in brain and liver. In turn, liver disease disrupts thiamine homeostasis and causes astrocytic damage. The latter results in the loss of neuron–astrocyte trafficking of neuroactive amino acids and thiamine

esters essential to CNS function. Thus, there are several possible routes to the production of brain damage in alcoholics. Further, these appear to be so interlinked that it may be unproductive to try and dissect the separate contribution of alcohol and thiamine deficiency to brain damage. Rather, it appears that a combination of malnutrition and alcohol intake can give rise to a range of cognitive deficits, from mild cognitive impairment to severe dementia, depending on the severity of the contributory abnormal mechanisms described by Butterworth.

Although the evidence points to alcohol being neurotoxic, the relationship between this and cognitive impairment is not clear. A major drawback of neuropathological studies is that such inferences have been either not possible or gleaned by retrospective case note analysis. In contrast, neuroimaging allows the prospective analysis of clinical features in relation to *in vivo* brain structural and functional abnormalities. Well-controlled, prospective studies using both CT and MRI have reliably confirmed the presence of cerebral shrinkage in alcoholics in life, involving both cortical grey and subcortical white matter²⁷⁻³⁴ which are more pronounced in older patients³⁵ and more apparent in the frontal lobe^{36,37}. The percentage of alcoholics with evidence of cerebral atrophy on brain scans is far in excess of that noted in neuropathological studies¹². This discrepancy is probably explained by the finding that brain changes are reversible with continuing abstinence^{29,33,34,38-42} indicating that neuroimaging findings do not accurately reflect the permanent neuropathological changes previously described. Some studies have found that the neuroimaging changes worsen in patients who continue to drink^{41,43,44}. A recent carefully controlled study, in which alcoholic men and controls were followed over 5 years⁴⁵ found that a greater total alcohol consumption was associated with greater decrease in cortical grey matter, particularly in the frontal lobe. Thus, it is possible that prolonged alcohol ingestion leads to irreversible brain damage, which might be mirrored by irreversible cognitive deficits.

There is certainly no doubt that cognitive dysfunction can be witnessed in uncomplicated alcoholics, even though these may not reach the severity required for a diagnosis of dementia. Parsons⁴⁶, following decades of his own research, concludes that both male and female sober adult alcoholics have deficits on tests of learning, memory, abstracting, problem-solving, perceptual analysis and synthesis, and speed of information processing, which are equivalent to those found in patients with known brain dysfunction of a mild to moderate nature. Such abnormalities can also be witnessed in adolescent alcoholics⁴⁷. Further attempts to identify factors other than alcoholism to account for these differences have been unsuccessful⁴⁶. It is noticeable, however, that, like neuroimaging studies, these deficits are largely reversible over weeks or years of abstinence^{46,48}. This may explain why studies relating cognitive and structural changes in alcoholics found weak and inconsistent correlations, the majority of which were explained mainly by age and premorbid IQ^{27-30,50}.

Functional imaging studies have been a bit more fruitful in this regard. One study found a reduction in overall cerebral glucose utilization⁵¹ but all other studies found specific regional changes. For example, Samson *et al.*⁵² have shown a relative decrease in glucose utilization within the medial frontal cortex of six recently detoxified neurologically intact chronic alcoholics, and Gilman *et al.*⁵³ have also reported the same finding in 14 alcoholics studied after at least 27 days of abstinence. In the latter study, patients made errors on the category sorting test, a test of frontal lobe function, and this correlated with glucose utilization in the medial frontal cortex. However, in a small-scale follow-up study from the same group, these deficits were partially reversible with abstinence⁵⁴ and it is not clear whether they become irreversible with continued drinking.

DISCUSSION

There are several conclusions that can be made from these studies concerning the validity of alcoholic dementia. First, diffuse cognitive impairment in alcoholic patients is common. It is an age-related phenomenon and the cognitive deficits are more mild than in patients with neurodegenerative dementias. Alcohol itself can produce irreversible brain damage, mainly of white matter and the frontal association cortex, but most studies suggest that frank dementia cannot be ascribed entirely to alcohol neurotoxicity. Other causes should therefore be sought, most notably thiamine deficiency. In younger patients, alcohol alone can also produce diffuse cognitive and cerebral abnormalities but these are largely reversible with abstinence. It remains to be determined which factors lead to the permanent alcohol-induced neuropathology seen at post mortem.

OTHER TOXIC DEMENTIAS IN THE ELDERLY

The acute and chronic psychiatric effects of ingestion of drugs of abuse, including those most relevant to the elderly—barbiturates and benzodiazepines—have been extensively reviewed by Lishman⁵⁵. Suffice it to say that there is little evidence to suggest that these produce diffuse cognitive impairment. The neuropsychiatric effects of long-term occupational solvent exposure is also relevant to the elderly. Although there is controversy concerning whether solvents *per se* produce long-lasting effects, most studies find evidence for lasting neuropsychological abnormalities⁵⁶⁻⁵⁹. Of particular interest is that these studies also found that a combination of long-term solvent exposure and alcohol abuse is a particular risk factor for the development of dementia. Of final relevance to this age group is the identification of bismuth encephalopathy, which results from the over-ingestion of bismuth-containing compounds commonly taken for gastric irritation and sold without prescription. Following an acute organic reaction characterized by delirium, seizures and neurological abnormalities, persistent sequelae of diffuse cognitive impairment and cerebral atrophy have been documented^{60,61}.

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Reversible Dementias

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The early literature on this subject includes numerous case series, or even single case studies, of “dementia” that seemingly resolved following treatment of the associated physical or psychiatric condition. Undoubtedly, many of the patients were actually suffering from delirium or other organic brain syndromes, an observation that is discussed in more detail by Byrne¹. Confusion over terminology has complicated research in this area. For example, the concept of “reversible” dementia appears at odds with the European use of the term “dementia”, denoting an insidiously progressive disorder². In addition, the relationship between delirium and dementia is a complex one and is dealt with elsewhere in this textbook.

This chapter reviews the classification and prevalence of *potentially* reversible dementias in clinical practice, the issue of response to treatment and the cost–benefit of routine investigation. A detailed presentation of the numerous causes of reversible organic brain syndromes is provided by Lishman².

CLASSIFICATION

A useful scheme, which clarifies terminology, has been suggested by Maletta³ and is shown in Table 53.1. Secondary dementias are those arising from a specific physical disorder. Drug intoxication and metabolic disorders include more acute conditions, often in association with impairment of consciousness, which clearly overlap with delirium but which have traditionally been included in studies of reversible dementia. Dementia due to psychiatric disorders includes those conditions classically referred to as the “pseudodementias”⁴ mostly secondary to primary depressive illness.

PREVALENCE

The frequency with which potentially reversible dementias occur in clinical practice is an important factor in determining the rigour with which contributory physical disorders or other causes are pursued. A previous review⁵ concluded that approximately 12% of patients presenting to a variety of specialist services with symptoms of dementia had treatable causes. This review was based on case series published during the 1970s and 1980s^{6–11}. The prevalence of potentially reversible disorders was 18% in patients under the age of 65 years but only 5% in those over 65. Depression, drug toxicity and normal pressure hydrocephalus accounted for more than half the cases. Identification rates were higher in specialist inpatient units than in outpatient or community-based studies. Importantly, though, these studies largely failed to review patients at a later date in order to determine the true reversibility of their cognitive symptoms.

Table 53.2 shows a selection of surveys of the potentially reversible dementias published during the 1990s^{12–18} grouped by specialty. Each was carried out in the setting of an outpatient memory clinic. Rates of identification of physical disorders were generally lower than in the earlier series. It has been suggested that the increasing awareness of the features of dementia and the use of internationally agreed diagnostic criteria has helped to refine clinical diagnosis¹⁹. This has resulted in a more rigorous exclusion of doubtful (non-dementia) cases at an early stage. Esoteric aetiologies are conspicuous by their absence, despite the comprehensive investigation carried out. This is likely to reflect the more representative nature of the patients studied. However, memory clinics specializing in the assessment of patients under the age of 65 usually still report higher rates of identification of potentially reversible conditions, particularly depression^{20,21}.

Table 53.1 Classification of potentially reversible dementias

Reversible dementias	Common clinical examples
1. Secondary dementias	
<i>Associated with neurological disorders</i>	
Structural lesions	Normal pressure hydrocephalus, brain tumour, subdural haematoma
<i>Associated with systemic disorders</i>	
Nutritional disorders	Vitamin B ₁₂ deficiency, folate deficiency
Endocrine disorders	Hypothyroidism, hyperthyroidism, hypoparathyroidism
Collagen/vascular disorders	Systemic lupus erythematosus, cerebral vasculitis
Infectious diseases	Neurosyphilis, chronic meningitis, AIDS
Alcohol related disorders	Primary alcoholic dementia
Miscellaneous	Chronic obstructive airways disease, sleep apnoea syndrome
2. Drug intoxication and metabolic disorders	Intoxication with major and minor tranquillizers, anti-hypertensives
3. Dementia due to psychiatric disorders	Depression, late-onset schizophrenia

After Maletta³.

Table 53.2 Outcome of investigation of patients presenting with clinical symptoms of dementia

Studies	Brody ¹² Almeida <i>et al.</i> ¹³	Ames <i>et al.</i> ¹⁴ Freter <i>et al.</i> ¹⁵	Chui & Zhang ¹⁶ Walstra <i>et al.</i> ¹⁷ Hogh <i>et al.</i> ¹⁸
Memory clinic specialty	Geriatric psychiatry	Geriatrics	Neurology
Total patients studied (<i>n</i>)	558	405	719
Clinical dementia confirmed [<i>n</i> (%)]	368 (100)	270 (100)	451 (100)
Mean age (range) (years)	70 (44–90)	76 (49–92)	66 (19–97)
Potentially reversible dementias			
<i>Secondary dementias</i> [<i>n</i> (%)]:	7 (1.9)	25 (9.3)	61 (13.5)
Normal pressure hydrocephalus	–	6	20
Brain tumour	–	2	1
Vitamin B ₁₂ deficiency	–	11	27
Folate deficiency	1	–	–
Hypothyroidism	1	3	8
Hyperthyroidism	–	–	1
Positive syphilis serology	1	3	4
Alcoholic dementia	4	–	–
<i>Drug intoxication and metabolic disorders</i> [<i>n</i> (%)]:	0	13 (4.8)	0
Alcohol abuse	–	2	–
Drug toxicity	–	11	–
<i>Psychiatric disorders</i> [<i>n</i> (%)]:	0	23 (8.5)	7 (1.6)
Depression	–	23	7
Follow-up period (months)	>6	>4	6 [†]
'True' reversible dementia [<i>n</i> (%)]	0	5 (1.9)	2 (0.4) [†]

[†]Not stated in Hogh *et al.*¹⁸

RESPONSE TO TREATMENT

The degree to which reversible dementias are, in fact, reversible has long been the subject of debate. Rabins²² reported some improvement in two-thirds of patients and complete recovery in 40%. This looks optimistic, judging by the results presented in Table 53.2 of between 0% and 2% reversibility.

A number of conditions deserve special mention. Depressive "pseudodementia" has traditionally been viewed as a treatable condition, with a distinct clinical history and symptoms that distinguish it from "true" dementia²³. In an attempt to improve the clinical discrimination between depressive pseudodementia and progressive dementia, Yousef *et al.*²⁴ derived a rating scale from a large number of possible discriminating features. Validating the diagnosis 12–14 months later, the scale allowed correct classification of 98% of true dementia cases and 95% of depression cases. Longer-term follow-up studies tend to support the view that severe cognitive impairment in depression is a harbinger of true dementia in 25–50% of patients^{24–26}. Alexopoulos *et al.*²⁶ found that, after 2–3 years, elderly depressed patients who also fulfilled DSM–III criteria for dementia were nearly five times more likely to develop dementia than those without cognitive impairment at presentation. The issue is further complicated by the high prevalence of depressive symptoms in dementia²⁷.

Several authors have questioned the usefulness of routine syphilis serology^{28–30}, while others have defended routine testing on the grounds that even one missed case would be catastrophic for the individual patient³¹. However, the erroneous diagnosis of active syphilis perhaps carries with it equally dire consequences, particularly when false positives may occur as a result of yaws and other non-venereal treponemal infection³². Hilton³³ has suggested that treatment should not be based solely on positive serology and that testing for syphilis should be dealt with in the same way as HIV testing and require informed consent where possible.

Normal pressure hydrocephalus is the most common neurological cause of reversible dementia^{16–18}. However, the results of shunting operations indicate high rates of post-operative

complications, including death, as well as a lack of evidence of effectiveness^{34–35}. The best results are obtained in patients under the age of 60 years who present with dementia of less than 6 months' duration³⁴.

Clearly, the lack of reversibility revealed by the studies listed in Table 53.2 suggests that most of the associated disorders are really *concomitant* with true dementia, rather than of aetiological significance. They may also be secondary to the dementia, e.g. anaemia as a result of malnutrition. Hogh *et al.*¹⁸ reported that 45% of those with dementia were found to have concomitant disorders. Although treatment of depressive symptoms in dementia has been shown to improve some aspects of cognition and everyday function^{36,37}, there is little evidence to show that the correction of other disorders, such as hypothyroidism or vitamin B₁₂ deficiency, has similar benefits^{17,38,39}.

Table 53.3 Physical investigations for the assessment of an elderly patient presenting with cognitive impairment

Routine investigations (all cases)	Special investigations (atypical cases or clinically indicated)
Full blood count ^{†a,b,c}	Brain CT or MRI ^{a,b,c}
Erythrocyte sedimentation rate ^b	EEG ^{a,c}
Renal function ^{a,b,c}	Chest X-ray ^{a,b,c}
Liver function ^{a,b,c}	Electrocardiogram ^{b,c}
Thyroid function ^{a,b,c}	Lumbar puncture ^{a,c}
Calcium ^{a,b,c}	
Vitamin B ₁₂ ^c	
Vitamin B ₁₂ ^{a,b,c}	Urine for culture ^{a,b}
Folate ^{b,c}	
Glucose ^{a,b,c}	HIV testing ^a
Syphilis serology ^{a,c}	Toxicity screen ^a
	Auto-antibody screen ^b

Recommendations for investigations: ^aAmerican Academy of Neurology⁴⁰; ^bRoyal College of Psychiatrists⁴²; ^cDutch Consensus statement⁴⁴.

INVESTIGATION PROTOCOLS

A number of organizations have published guidelines for the investigation of dementia, largely with the aim of identifying the common reversible dementias⁴⁰⁻⁴⁴. Table 53.3 lists investigations suggested in three sets of consensus statements^{40,42,44} and shows that there is broad agreement on the appropriate routine blood investigations. Chui and Zhang¹⁶ examined the added value of the investigations recommended by the American Academy of Neurology⁴⁰. After a complete clinical assessment, blood tests and neuroimaging results changed management in only 13% and 15% of cases, respectively. van Crevel *et al.*⁴⁴ found that the number of patients requiring investigation to find one case of reversible dementia was approximately 100, and that any financial saving on care costs was insignificant. Lastly, Foster *et al.*⁴⁵, in a detailed examination of the cost-effectiveness of routine computed tomography (CT), concluded that patients aged over 65 years should only be scanned if symptom duration was less than 1 year, if symptoms were rapidly progressive or if the presentation was atypical of Alzheimer's disease. Routine scanning was recommended in all patients presenting under the age of 65.

CONCLUSION

Reversible dementias are rare but their identification is important from the individual patient's perspective. The literature encompasses a polarization of views. The evidence-based view would suggest that extensive investigation is unnecessary and that, in any case, treatment of identified disorders is largely ineffective. The more traditional, patient-centred view, perhaps coloured by the early literature and the fear of litigation, would demand the continued use of routine investigations in the hope of excluding treatable aetiologies, however rare. The introduction of care protocols reached by consensus should guide clinicians and ensure that patients have equity of access to appropriate assessment.

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Differential Diagnosis of Dementia

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When approaching the differential diagnosis of dementia, two distinct but related issues are involved. First, the clinician has to consider whether the patient is suffering from a dementia syndrome. Differentiation has to be made from a functional psychiatric disorder, such as depression, which may be manifested as a dementing illness (so-called "pseudodementia" or the dementia syndrome of depression); from an acute organic reaction (acute or subacute confusional state); from the effects of normal ageing¹, from pre-existing handicaps; and from the deleterious effects of drugs. Second, the aetiology of the dementia, in terms of either a potentially reversible or irreversible dementia, has to be uncovered. The process is illustrated in Figure 54.1, although in practice a clinician formulates a diagnosis based on the whole presentation (see also ref. 2). This chapter will not deal with Alzheimer's disease in detail (see ref. 3) but it will be regarded as a prototype against which other disorders can be compared.

There are clinical situations in which the diagnosis of dementia is problematic. These have been well summarized⁴ and include: early cases (where the effects of normal ageing need to be considered); patients with a low IQ (where intellectual symptoms may be noticed early); very old patients, especially when residents of nursing homes, without reliable informants; patients with impaired vision and hearing; patients who are mentally handicapped; patients with prominent psychiatric problems, such as paranoia, dysthymia or personality problems; and those with severe physical illness or an intercurrent delirium.

When evaluating a patient with possible dementia, the following should be performed⁵⁻⁷:

- Detailed family and personal history and history of the current illness from a reliable informant.
- Mental State Examination of the patient, with particular reference to the cognitive state (amnesia, apraxia, aphasia and agnosia); more detailed neuropsychological assessment should be considered if particular deficits are suspected).
- Physical examination, with particular emphasis on the central nervous system.
- Investigations including haematological and biochemical blood tests, serum B₁₂ and folate, thyroid function tests, chest X-ray and ECG.

Some form of assessment of cerebral function/structure is desirable, the nature of the investigation being dependent on local facilities. An electroencephalogram (EEG) should be performed and is widely available. Ideally, a computed tomography (CT) scan should also be carried out, but this is not always practicable. If so, it is reasonable to limit this examination to cases

in whom there is reason to suspect an intracranial lesion, i.e. clinical suspicion of such a lesion, evidence of cerebral infarction, focal neurological signs, seizure activity, a head injury thought to be contributory to the clinical picture, or a suspicion of normal pressure hydrocephalus. The main conditions in the differential diagnosis of dementia and possible causes of the dementia syndrome are discussed below.

ACUTE CONFUSIONAL STATE (DELIRIUM)

This is the most important differential diagnosis to be considered, as there is almost always a physical disorder underlying its presence. Onset is acute, disturbances often severe and the patient is usually brought to the attention of the services by worried friends, relatives or neighbours. There is a global disturbance of cognition, with marked fluctuation over the course of the day, often worse at night⁸.

During affected periods, there is almost always some disorientation. Among the disturbances are disorders of perception (characterized by an inability to interpret events and to discriminate them from images and dreams), disorders of thinking (disorganized, fragmented and with disjointed thoughts and decreased ability to plan or solve problems) and disorders of memory (registration, retention and retrieval are all affected). Clouding of consciousness is the cardinal feature of delirium and has been defined in terms of a disorder of attention (decreased or increased alertness, selectiveness and directiveness) and wakefulness (diminished night sleep⁹). Psychomotor behaviour is usually disturbed, with overactivity, underactivity or a mixture of the two. The diagnosis of acute confusion is made on a characteristic history of a sudden onset of disturbance and the findings outlined above on examination of the patient. Often the physical precipitant is not obvious initially (and in a proportion of cases is never discovered). An underlying dementia may be present and is suggested by the history.

PSEUDODEMENTIA

In this condition, symptoms of disorientation and memory loss occur in a non-organic psychosis and mimic dementia. The original description of cases included patients with a number of functional psychiatric disorders¹⁰, but in practice depression is by far the commonest cause. In contrast to dementia, the clinical course of the condition is relatively short and has a defined date of onset. There is often a previous history and/or a

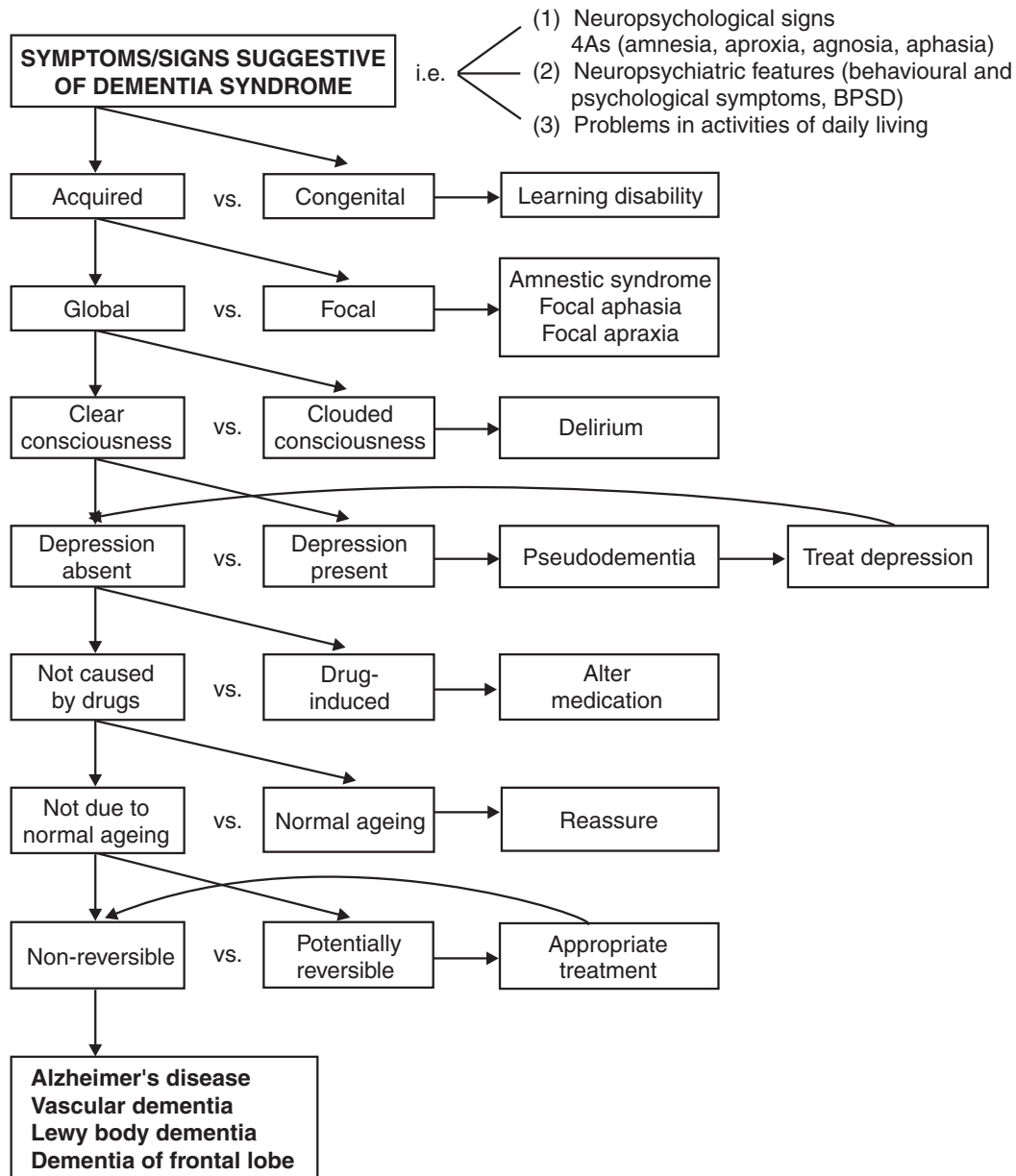


Figure 54.1 Diagnostic algorithm for dementia

family history of affective disorder. Patients tend to answer questions with “don’t know” responses, there is a variability in their ability to perform tasks of similar difficulty, they communicate a sense of distress and complain of their cognitive deficits¹¹. Structural brain changes have been described in this condition¹². Pseudodementia is currently a term in disrepute and the alternative (dementia syndrome of depression) is considered to be more accurate. The diagnosis of pseudodementia is made on clinical grounds and should always be considered in a demented patient. Validation of the diagnosis is by successful treatment and return to normal of the cognitive deficit^{13,14}. Some clinicians would argue that a trial of antidepressants should be given in all cases of dementia.

VASCULAR DEMENTIA

There are two main types of vascular dementia, one involving subcortical structures affecting small arteries, leading to the clinical picture of subcortical dementia (see below for cardinal features), and one involving medium-sized arteries (anterior, middle and posterior cerebral arteries), leading to a cortical dementia. The clinical features of vascular cortical dementia have been defined in the Hachinski score¹⁵. Physical examination will generally reveal neurological signs, such as disturbances of gait rigidity, spasticity and reflex abnormalities, and CT often reveals cerebral infarction. Diagnostic criteria have been described^{16,17}.

INTRACRANIAL LESIONS

Any form of intracranial mass lesion may masquerade as dementia. Cerebral tumours (primary or secondary), brain abscess or intracranial bleeds may all give rise to a dementia syndrome. Diagnosis is made through suspicion of the primary lesion and visualization on CT scan.

SUBCORTICAL DEMENTIA

This is a generic term for a particular syndrome of dementia, which has particular clinical features separating it from a cortical dementia. These include mental slowness, inertia, apathy and loss of initiative, occurring along with cognitive disturbances. The diseases associated with subcortical dementias include Parkinson's disease, Huntington's disease, progressive supranuclear palsy, Wilson's disease, spinocerebellar degeneration, hydrocephalus and the toxic/metabolic encephalopathies. The structures affected are subcortical—the basal ganglia, thalamus and the brain stem.

In Parkinson's disease, there is mental slowing, diminished problem solving, poor memory and a deterioration in abstraction, concept formation and word generation. Depression is common, as in all subcortical dementias. Cortical features (such as aphasia, apraxia and agnosia) are absent, although some authors claim there is a cortical dementia present, probably due to coexisting Alzheimer's disease. The diagnosis of Parkinson's disease is essentially a clinical one and physical manifestations of the disease will generally be present when the dementia is apparent.

Huntington's disease is characterized by choreiform movements and associated with autosomal dominant transmission. Personality changes, irritability and apathy are the first changes and predate the chorea. The dementia appears soon after the movement disorder becomes apparent, characterized by memory disturbance (impaired recall of both recent and remote memories), slowing, failure to initiate cognitions (especially those required in planning) and impaired concentration and judgement²⁶. Although dementia without chorea has been described, it is very rare and the diagnosis of Huntington's disease is usually suspected prior to the onset of dementia¹⁸.

Pseudobulbar palsy, rigidity (more pronounced in the neck and trunk) and paralysis of vertical gaze (downward gaze lost first, followed by failure of upward gaze) are hallmarks of progressive supranuclear palsy, and the associated dementia is classically subcortical (indeed, the original description of subcortical dementia was based on the dementia of progressive supranuclear palsy). Speech is disrupted (e.g. dysarthria and hypophonia) but aphasia is absent. The clinical features are such that the diagnosis will often be suspected and, although it may be confused with Parkinson's disease, the characteristic tremor of the latter is absent.

Other conditions resulting in subcortical dementias occur rarely and are of limited relevance to old age psychiatry. The amyotrophic lateral sclerosis–Parkinson–dementia complex of Guam is rare outside the Chonorro population of the Western Pacific, the dementia being profound and characterized by features of decreased memory, apathy and slowness; cerebellar degenerations are associated features of subcortical dementia but the cerebellar dysfunction is usually the prominent feature; Wilson's disease and Friedreich's ataxia are associated with dementia, but are confined to children and young adults.

NORMAL PRESSURE HYDROCEPHALUS

A potentially treatable cause for a dementia syndrome is hydrocephalus, the most widely cited being normal pressure hydrocephalus. The classical clinical triad consists of gait disturbance (ranging from mild clumsiness to akinesia), incontinence (almost always urinary incontinence, occurring late in the illness) and dementia (impaired memory, disorientation and mental slowing). The diagnosis is made by radiological examination (CT scan shows marked enlargement of the ventricles with relatively normal cortical sulci, and isotope cisternography shows obstruction to the flow over the cortex). A ventricular shunt to divert cerebrospinal fluid is the appropriate treatment but not all cases improve, even when the classical clinical picture is present.

PICK'S DISEASE

Personality changes and mood disorders (ranging from depression to elation) occur first, with coarsening of affect and antisocial behaviour. Impaired judgement occurs, with loss of insight. Aphasia and circumlocution occur early. Memory is relatively unimpaired until the later stages of the disease, as is praxic function. The Kluver–Bucy syndrome has been described early in the illness, but this also appears in Alzheimer's disease¹⁹. Extrapyramidal signs appear late in the illness but the clinical picture is unlikely to be confused with Parkinson's disease. Seizures are said to be less common than in Alzheimer's disease and CT scan shows frontal and temporal lobe atrophy, rather than the generalized shrinkage seen in Alzheimer's disease. The diagnosis is made on the basis of the characteristic onset of personality changes before the onset of dementia.

DEMENTIA OF THE FRONTAL LOBE TYPE

Recently, descriptions have emerged of a form of dementia that appears to affect the frontal lobes and is defined in terms of clinical presentation (personality change, speech impairment and relative preservation of visuospatial functions), neuropsychological function (frontal lobe syndrome) and blood flow studies (diminished frontal lobe blood flow on single-photon emission tomography)^{20–22}. There is frontal and temporal lobe atrophy and, while the condition resembles Pick's disease, the characteristic neuronal inclusion bodies are absent. Diagnosis is made on clinical grounds but confirmation usually has to await autopsy.

DEMENTIA OF LEWY BODY TYPE

This has been described with increasing frequency and is considered to be amongst the commonest forms of primary dementia²³. The relationship of this syndrome to Parkinson's disease is uncertain, but the clinical picture of parkinsonian features, hallucinosis and episodes of confusion should raise the diagnostic possibility.

TOXIC-METABOLIC ABNORMALITIES

Generally, these are readily identifiable by the primary cause for the syndrome (such as anoxia, renal or hepatic failure) and the symptomatology produced is more often identifiable as an encephalopathy rather than a dementia syndrome. There are a

few conditions of relevance in old age psychiatry, which will be outlined briefly. Systemic carcinomas may produce effects in the brain by several mechanisms through metabolic disturbances (such as excess secretion of adrenocorticotrophic hormone or antidiuretic hormone), through structural change (cerebral secondaries or infections) or through remote effect (limbic encephalitis). This last condition is common in men, particularly with oat cell carcinomas of the lung, and has a course of up to 24 months. Affective changes dominate the picture, with amnesia the primary (and occasionally the sole) abnormality. Diagnosis of the condition is through attention to the physical condition of the individual. Vitamin deficiencies (notably B₁₂ and folate) cause mental impairment, but routine testing for their levels makes the diagnosis relatively easy. The same holds true for thyroid dysfunction and hypercalcaemia. Chronic excessive alcohol intake can result in a dementia²⁴.

CREUTZFELDT–JAKOB DISEASE

Early features include fatigue and listlessness, elevated mood and impaired memory and concentration. Motor abnormalities occur with spasticity, ataxia and tremor. Myoclonic jerks and seizures may occur. The EEG is very abnormal, with characteristic triphasic waves superimposed on some suppression of the background rhythms. The course of the illness is very rapid and most affected individuals are dead within 2 years²⁵.

SUMMARY

The differential diagnosis of dementia is a two-stage process: first, the differentiation of dementia from other causes of cognitive impairment; and second, if it is found to be a form of dementia, the elucidation of the aetiology. Alzheimer's disease is the commonest form of dementia, cerebrovascular disease is probably the second commonest cause and Lewy body dementia is becoming increasingly recognized. The differential diagnosis of dementia is an excellent example of how simple logical clinical skills can be applied without the need for expensive investigations, which should be reserved for situations where there is clinical doubt about the diagnosis.

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Distinguishing Depression from Dementia

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Recent research continues to confirm the complex relationship between depression and dementia. The ability to distinguish depression from dementia is complicated by the overlap of many of their clinical manifestations. Differentiating the two illnesses is further complicated in that depression and dementia are often concurrent illnesses in the geriatric population. Depression has also become recognized as a possible prodrome, or even risk factor, for degenerative dementias. Finally, cerebrovascular disease, one of the long-established etiologies of dementia, is becoming increasingly recognized as a possible etiology of geriatric depression.

Dementia and depression are two of the most common diagnoses that clinicians encounter in geriatric psychiatry. The prevalence of dementia has been estimated to be approximately 1% at age 60, doubling every 5 years to reach 30% to 50% by age 85^{1,2}. Major depression afflicts 1–2% of community-dwelling elderly, with significantly higher rates observed in hospitalized elderly and those residing in nursing homes³. The prevalence for minor depression or subsyndromal depression is even higher, with rates reported to be 13–27%³.

Given the complex relationship between depression and dementia, combined with high prevalence rates, clinicians need to be familiar with the literature that addresses these issues. This chapter will begin with a review of the basic diagnostic criteria of dementia and depression. It will then explore the key points in clinically distinguishing the diagnoses. Next, it will examine the issue of concurrent depression and dementia, as well as the concept of pseudodementia of depression. It will conclude with a discussion regarding the latest research on the possibility of depression as a herald or prodrome of degenerative dementias, and the newly emerging concept of vascular depression.

CLINICAL PRESENTATIONS OF DEMENTIA AND GERIATRIC DEPRESSION

The DSM-IV diagnostic criteria for dementia include the development of multiple cognitive deficits manifested by memory impairment, and one or more of the following: aphasia, apraxia, agnosia or disturbance in executive functioning⁴. In addition to these core cognitive symptoms, numerous psychiatric symptoms are also common. In Alzheimer's disease (AD), the most common of the dementing disorders⁵, significant psychiatric symptoms often occur, including personality changes, irritability, anxiety, delusions, hallucinations and depressive symptoms^{5–8}. The depressive symptoms that are common in dementia include sleep disturbance, anorexia, irritability, social withdrawal, anergy and apathy^{6,7}.

The diagnostic criteria for major depressive disorder, as described in the DSM-IV, include depressed mood, diminished interest, weight loss, sleep disturbance, psychomotor agitation or retardation, loss of energy, feelings of worthlessness or guilt, and diminished ability to concentrate⁴. While not all studies agree, most authors report that in the elderly, compared to younger depressive counterparts, depressed mood and feelings of guilt are not as common but somatic and cognitive symptoms are more common^{3,9}. The cognitive symptoms may become so severe as to lead to the development of what has been termed "pseudodementia".

With such significant overlap of the symptoms of dementia and depression in the elderly, the clinical or "bedside" differentiation can be quite challenging. Reynolds *et al.*¹⁰ found that patients with pseudodementia of depression showed greater early morning awakening, higher anxiety and more severe impairment of libido. Patients with dementia showed more disorientation to time and greater difficulty with dressing and navigating through familiar surroundings.

The American Association for Geriatric Psychiatry, the Alzheimer's Association and the American Geriatric Society presented a consensus statement that included tips for differentiating dementia from depression⁵. The panel reported that patients with AD, in comparison with depressed patients, tend to minimize cognitive deficits, demonstrate impaired memory and executive function, have "indirect" symptoms of depression, such as agitation and insomnia, and demonstrate other cognitive deficits, such as aphasia and apraxia. The panel further reported that those patients with cognitive disturbance, in the context of depression, in comparison with demented patients, tended to exaggerate cognitive deficits, show impaired motivation and classic mood symptoms and have intact language and motor skills.

Abram and Alexopoulos¹¹ described the similarities between the clinical appearances of dementia and depression, including shared neurovegetative signs, such as weight loss, insomnia, decreased libido and fatigue. However, they also emphasized that some distinctions can be made clinically. Specifically, the authors asserted that weight loss, fatigability and insomnia usually reflect acute changes in depression, but will be more chronic in dementia. Mood-incongruent delusions may be found in psychotic depression as well as dementia, but mood-congruent delusions are more characteristic of depression. They also reported that the agitation of dementia often manifests as "pacing", while "hand-wringing" is more characteristic of depression.

Other features of the patient's presentation, as well as personal and family history, can be helpful in distinguishing depression from dementia. Geldmacher and Whitehouse² suggested that, in patients with dementia, more often a relative will report decreased

memory in the patient. In contrast, in patients with depression, if cognitive symptoms are reported, it is usually by the patient. They also stated that the duration of the presentation could be helpful in clarifying the diagnosis. Depression is typically of shorter duration than dementia and usually with a more well-defined onset. The authors also recommend a review of personal or family history for depression, and a review of family history for dementia.

SCREENING INSTRUMENTS

The use of various “bedside” screening instruments has been proposed as a way to clarify the diagnoses of dementia and depression. Common instruments used in the diagnosis of depression include the Hamilton Rating Scale for Depression (HAM-D), the Beck Depression Inventory (BDI), and the Montgomery–Asberg Depression Rating Scale (MADRS). In general, many authors have found that general depression screening instruments are less helpful in the diagnosis of depression in the elderly than in the general adult population. Reasons for this age disparity include decreased specificity, due to overlap of many depression symptoms with other common geriatric presentations, including dementia and general medical illnesses. It is also believed that the sensitivity is decreased, as not all of the classic symptoms of depression are as common in the geriatric population as in the general adult population.

A number of depression screening tools specifically designed to be used in geriatric or demented patients have been developed. Among the more commonly used instruments are the Geriatric Depression Scale (GDS), Alzheimer’s Disease Assessment Scale (ADAS) and the Cornell Scale for Depression in Dementia (CS). The ADAS relies on observation of the patients and thus avoids many of the difficulties inherent in interviewing a cognitively impaired individual¹¹. The CS combines interviews with the patient and caregivers, focusing on behavior during the week preceding the interview, thus taking advantage of available collateral information that many other scales ignore^{11,12}.

The most commonly used dementia screening tool is the Mini-Mental State Examination (MMSE). The MMSE was developed by Folstein *et al.* in 1975¹³. Since that time, it has become the most widely used examination for rapidly assessing the cognitive status of the elderly¹⁴. This short test covers a broad range of cognitive domains, including orientation, registration, attention and recall, calculation, language, and constructional ability¹⁵. Age- and education-specific reference values have been developed to help guide the clinician in the use of the MMSE in various populations¹⁴.

Other tests that have been developed to screen for cognitive impairment in the elderly include the Blessed Dementia Rating Scale and the Short Portable Mental Status Questionnaire (SPMSQ). A quick and simple-to-administer test, the clock-drawing test, has been shown in numerous studies to be a good screening test for dementia¹⁶. Numerous methods have been developed to score the clock-drawing test, ranging from complex 20-point scales to more simple ordinal scales.

Watson *et al.*¹⁷, developed a scoring system that is based on a seven-point scale. With this method, the patient is instructed to draw numbers within a pre-drawn circle to make the face of a clock. After completion, the clock face is divided into quadrants and the number of digits in each quadrant is counted. An error score of one is assigned for each of the first three quadrants containing any erroneous number of digits and an error score of four is assigned for the fourth quadrant if it contains an erroneous number of digits. A score of 4 or greater has been shown to have a sensitivity of 87% and a specificity of 82% in screening for dementia¹⁶.

Kafonek *et al.*¹⁸ conducted a study to determine the sensitivity and specificity of the MMSE in detecting dementia and the Geriatric Depression Scale (GDS) for detecting depression in an academic center-affiliated nursing home population. They found that, in screening for dementia, the sensitivity of the MMSE was 81% and the specificity was 83% when using a cut-off score of 24/30. The GDS was found to be less sensitive and less specific in screening for depression, with a sensitivity of 47% and a specificity of 75%. Thus, while such screening tools can provide useful additional information, they should be taken into consideration as part of the overall clinical picture and not used as the sole basis for making or excluding a diagnosis.

NEUROPSYCHOLOGICAL ASSESSMENT

Formal neuropsychological testing is often employed to help diagnose dementia and depression in the geriatric population. Neuropsychologists are able to administer a full battery of tests that have at least three distinct functions in the assessment of geriatric patients¹⁹. The first goal of neuropsychological testing is to aid with differential diagnoses between normal aging, psychiatric disorders and neurodegenerative/dementing disorders. A second common use of formal testing is to establish a baseline from which changes can be tracked over time. This can help in determining the response to treatment for depression or dementia, as well as to systematically follow the progression of dementing disorders. The final common use of neuropsychological testing is to delineate the strengths and weaknesses of a particular patient, to help make clinical recommendations for treatment, daily activities and planning for the future.

In an exhaustive review of the neuropsychological testing literature dealing with the differential diagnosis of major progressive dementias and depression, Rosenstein²⁰ summarized the major test variables that neuropsychologists have found helpful in distinguishing depression from various dementing disorders. In general, she reports that depressed patients tend to demonstrate normal to slightly reduced attention, memory, visuospatial functions, language, executive function, reasoning and sensory-motor function, with a negative or empty response style and inconsistent performances (even within the same domain). In contrast, patients with AD tend to demonstrate more significant impairment in memory, verbal fluency and executive function (in particular, poor awareness of deficits), with a higher prevalence of false-positive responses and intrusions.

Obviously, the milder the dementia at the time of testing, the milder will be the above-mentioned deficits. However, even mild deficits will be more accurately detected on formal neuropsychological testing than with the usual clinical assessment, or with screening tests such as the MMSE. This increased sensitivity is one of the strengths of formal neuropsychological testing. As part of a prospective epidemiological study of dementia, Jacobs *et al.*²¹ administered a comprehensive neuropsychological test battery to a group of initially non-demented older adults, in order to examine the association between baseline scores and subsequent development of dementia. The results of the study indicate that, even in the preclinical phase of AD, deficits can be found in areas of word-finding ability, abstract reasoning and memory.

MacKnight *et al.*²², in their paper examining the factors associated with inconsistent diagnosis of dementia between physicians and neuropsychologists, concluded that the two disciplines have complementary strengths. The neuropsychologist has superior skills in identifying early cognitive loss, while the physician’s expertise arises from a greater ability to assess the impact of the impairment on the patient’s ability to function. Thus, if available to the clinician, formal neuropsychological testing can provide very helpful and complementary information.

This is particularly true in those cases where diagnostic uncertainty is high.

LABORATORY EVALUATION

For many years attempts have been made to find a reliable biological marker for depression that could be used to develop a sensitive and specific laboratory test. Early on, investigators examined the use of the dexamethasone suppression test (DST), a neuroendocrine measure of the functioning of the hypothalamic–pituitary–adrenal axis, to distinguish depression from dementia. Although the test appeared to have some discriminatory ability in some of the very early studies^{23,24}, subsequent research failed to demonstrate its usefulness^{25–29}. Presently, the DST is not a routine part of a depression or dementia work-up.

Laboratory studies that should be a routine part of the assessment of dementia include a complete blood count (CBC), urinalysis, serum chemistry panel, liver function tests, thyroid function, serum vitamin B₁₂ and folate levels, and syphilis serology^{8,30}. A number of these tests are also useful in the evaluation of depression. The most commonly cited are a CBC, serum electrolytes, serum vitamin B₁₂ level, serum folate level, and thyroid function. While these laboratory studies do not necessarily help to distinguish depression from dementia, they can help to evaluate a number of possible common etiologies and contributing factors. Should any of these tests have positive results, steps should be taken to correct the abnormality and the patient should then be reassessed for depression or dementia.

Genetic testing is beginning to receive much attention as a possible aid in diagnosing various dementing disorders. Early-onset AD comprises both sporadic cases and those that are now known to represent a collection of single-gene disorders. Specific mutations in three genes have been associated with the early-onset, familial-pattern AD. The identified genes are the amyloid precursor protein (APP) gene on chromosome 21, presenilin-1 (PS1) gene on chromosome 14, and presenilin-2 (PS2) gene on chromosome 1³¹. In cases where early-onset dementia is suspected, genetic testing is indicated to help confirm the diagnosis and to provide additional information to be used in counseling the patient and family members.

In the more common late-onset form of AD, one gene, apolipoprotein E (APOE), located on chromosome 19, has been associated as a risk factor. This gene has three common isoforms, $\epsilon 2$, $\epsilon 3$ and $\epsilon 4$. Research has shown that the isoform $\epsilon 4$ is associated with the development of AD in a dose-dependent manner, and that the isoform $\epsilon 2$ may be protective for AD in a dose-dependent manner^{31–35}. Thus, the more copies of the $\epsilon 4$ allele one carries in one's genotype, the higher the risk for developing AD. One complicating factor in using this genetic test in the differential diagnosis of dementia and depression is that the $\epsilon 4$ allele has also been associated with the development of late-onset major depression in a study by Krishnan *et al.*³³ While genetic testing for APOE isoforms can be another helpful piece in distinguishing dementia and depression, it is far from conclusive and is currently not a routine part of a work-up for depressed, cognitively-impaired patients.

ELECTROENCEPHALOGRAPHY EVALUATION

A great deal of research has been conducted examining the ability of electroencephalography (EEG) studies to aid in the diagnoses of depression and dementia. In a literature review of EEG of the elderly, Klass and Brenner³⁶ reported several reasons why an EEG may be helpful in the evaluation of dementia. First, it may confirm that an abnormality of cerebral function exists. Second, it

may indicate that a focal process is present, rather than a diffuse process. Third, it may find that a previously unidentified seizure disorder is present. Finally, certain dementing processes, such as Creutzfeldt–Jakob disease, have pathognomonic EEG change and thus the EEG can be diagnostic in those cases. The authors concluded that, in depression, the awake EEG is usually normal and when abnormalities are found they most often are mild.

As abnormalities of sleep are part of the core symptoms of depression, sleep studies using EEG have been studied extensively for many years in an attempt to find objective variables that could aid in the diagnosis of depression. Variables that have consistently been found in depression include increased rapid eye movement (REM) activity, increased sleep latency, decreased REM latency, increased REM density and high rates of sleep onset REM as compared to normal control subjects^{37,38}. Dykierek *et al.*³⁸ studied the REM sleep parameters that could be useful in differentiating AD from normal aging and depression in the geriatric population. They found that they were able to correctly classify 86% of the patients studied using REM density and REM latency measures. In their study, depressed patients tended to have increased REM density and decreased REM latency, while demented patients tended to have decreased REM density and normal to increased REM latency. While the use of EEG and sleep studies are not a standard part of a dementia and depression work-up, in selected cases they may provide additional information that could be helpful to the clinician.

NEUROIMAGING

Recent years have seen significant advances in the development of technology for imaging the central nervous system. Although the value of neuroimaging in the work-up of dementia and geriatric depression is controversial, many authors present strong arguments for its diagnostic utility. The rationale for the use of magnetic resonance imaging/computed tomography (MRI/CT) scans in the work-up of dementia includes the following: (a) neuroimaging contributes to increased diagnostic accuracy and may detect occult lesions not evident on clinical examination; (b) most physicians' clinical skills in diagnosing AD are not sufficient to abandon imaging; (c) CT or MRI may identify potentially treatable causes of dementia missed by clinical evaluation; (d) imaging protects against possible malpractice suits³⁹. The arguments against the routine use of neuroimaging include: (a) lack of cost-effectiveness and low yield (prevalence of < 5% for clinically significant findings); (b) lack of influence on eventual outcome; (c) imaging may result in unwanted procedures (e.g. surgery) or cause distress to patients; (d) benign small vessel changes on MRI often result in overdiagnosis of vascular dementia (false positives)⁴⁰.

Historically, the use of neuroimaging in the work-up of depression has not been as strongly recommended as it has in the work-up of dementia. However, recent research of the possibility of vascular and degenerative etiologies of geriatric depression has raised the issue of whether neuroimaging should be part of the work-up of geriatric depression⁴¹. As will be discussed later in this chapter, geriatric depression is becoming increasingly recognized as a heterogeneous illness with multiple possible etiologies, including cerebrovascular disease and neurodegenerative disease, as well as the genetic and psychosocial etiologies recognized in the general adult population. As these possible etiologies become better established, the use of neuroimaging may become more helpful in the classification of geriatric depression, thus impacting treatment and prognosis.

Recent research on the diagnosis of AD using MRI estimated measurements of the temporal lobes has demonstrated impressive sensitivity and specificity. O'Brien *et al.*⁴², in a study to determine the utility of temporal lobe MRI in distinguishing AD from

depression, normal aging and other causes of cognitive impairment, found that temporal lobe atrophy provided “good separation” between those patients with AD and those with other causes of cognitive impairment. In particular, anterior hippocampal atrophy was associated with a sensitivity of 83% for detection of AD, and a specificity of 80% for controls, 87% for depressed subjects and 89% for others. Jack *et al.*⁴³ found that measurements of hippocampal atrophy provided a sensitive marker for AD, even in the early stages. However, recent studies linking smaller hippocampal volume with depression may limit the utility of this measure in distinguishing depression from dementia^{44,45}.

The use of functional neuroimaging technology, such as positron-emission tomography (PET) scans and single photon emission tomography (SPECT) scans in the evaluation of dementia and depression is increasingly becoming a subject of research. Although some interesting findings are beginning to emerge⁴⁶⁻⁴⁸, functional neuroimaging remains primarily a research tool, with limited application to clinical practice. As the technology becomes less expensive and the research shows more consistent and diagnostically useful findings, functional imaging studies may prove to be very powerful tools.

Overall, the use of any form of neuroimaging as a routine part of the work-up of dementia and geriatric depression remains controversial. The utility of functional neuroimaging is uncertain and its use is limited mainly to research. The utility of structural neuroimaging, such as CT or MRI, is becoming more accepted as part of a dementia work-up, but is just beginning to be considered a part of the work-up for geriatric depression. In distinguishing dementia from depression, these studies can provide another useful piece of information to the clinician. However, as with all diagnostic tests, they are not without expense and potential risks/harms, and should be utilized only when the remainder of the clinical evaluation warrants it.

CONCURRENT DEPRESSION AND DEMENTIA

To this point, depression and dementia have been discussed as though they were mutually exclusive diagnoses. However, they commonly coexist in the geriatric population. This concurrence further complicates the task of making a clear diagnosis in the patient who presents with a combination of mood symptoms and cognitive symptoms. Clinical studies report a wide range of prevalence rates for depression in dementia patients. This variance appears to depend on the criteria used and the population studied. In patients with AD, reported rates for depressive disorders have ranged from 0% to 86%, with most studies showing 30–40%^{12,49}. The reported rates of depressive symptoms that do not reach diagnostic criteria for a mood disorder are even higher. Many of the less common dementing disorders, such as Parkinson’s disease, Huntington’s disease and frontotemporal dementia, have rates of depression that exceed those found in AD^{50,51}.

In light of the significant co-morbidity of depression and the dementing disorders, a number of authors^{12,49} are now arguing that the clinical approach to evaluating the elderly patient with mood symptoms and cognitive symptoms should move from the traditional either/or approach to an and/or approach. The impact of not recognizing depression in the context of dementia is significant. Concurrent depression has been reported to increase the disability of patients by lowering the ability to perform instrumental activities of daily living (IADLs) in the mildly cognitively impaired, and lowering the ability to perform activities of daily living (ADLs) in the more severely cognitively impaired⁴⁹. Concurrent depression has even been associated with a greater than 59% mortality rate during the first year in elderly patients admitted to nursing homes⁴⁹. Thus, the recognition and treatment

of depression in the context of dementia can have a profound effect on morbidity and mortality in elderly patients.

“PSEUDODEMENTIA” AND DEPRESSION AS A PRODROME TO DEMENTIA

A great deal has been written about depression with cognitive impairment, generally referred to as “pseudodementia”. This has been classically listed as one of the most common causes of reversible dementia. The most commonly described clinical scenario would be of an elderly patient undergoing an evaluation for cognitive impairment and being found to meet diagnostic criteria for depression. With appropriate treatment and resolution of the depression, a significant proportion of such patients would no longer have identifiable cognitive impairment. Increasing evidence, however, has convincingly shown that depression with an initially reversible dementia syndrome will frequently be followed by the development of an irreversible dementia^{52,53}.

Recent studies have also shown that depressive symptoms in the geriatric population, even without evidence of “pseudodementia”, may be a prodrome of degenerative dementias. In a large epidemiological study, Berger *et al.*⁵⁴, found a significantly higher number of depressive symptoms in individuals who later developed AD than in those individuals who did not go on to develop AD. Reding *et al.*⁵⁵ found that, in a sample of patients evaluated in a dementia clinic, 57% of the depressed, non-demented patients develop frank dementia at a 3 year follow-up. The relationship between preclinical depression and the development of dementia is complicated by the difficulty in discerning whether the depression is truly part of a prodrome phase of dementia or whether it is a risk factor for later development of dementia⁵⁵. In either case, the former concept of pseudodementia, separate and apart from true dementia, is not as clear as it once seemed.

VASCULAR DEPRESSION

Recent research has indicated yet another factor that further complicates the relationship between dementia and geriatric depression. It has long been recognized that depression is a common complication of strokes. It is now beginning to be recognized that depression and mood symptoms can be secondary to “silent strokes” or the accumulation of microvascular lesions in the CNS^{41,56-58}. Numerous studies are now reporting that significant proportions of elderly patients with depression have evidence of cerebrovascular disease found on MRI and CT studies^{56,57}. In particular, lesions in the basal ganglia, periventricular white matter and deep cortical white matter have been associated with depressive symptoms and depression in the elderly^{59,60}.

This has led to the development of the “vascular depression” hypothesis. Like the dementia in vascular dementia, the depression in vascular depression is thought to arise from vascular insults to the central nervous system. Instead of affecting the systems involved in cognitive processes, the injuries are thought to affect the systems involved in the maintenance of a normal affective state. Thus, it is possible that cerebral vascular disease can not only result in classic stroke symptoms but can also lead to the development of dementia and/or depression.

CONCLUSION

Distinguishing depression from dementia is a difficult clinical task. The cornerstone of making an accurate diagnosis remains the

clinical information gathered "at the bedside" from the patient and his/her family. Various screening instruments and routine laboratory studies can contribute useful additional information to the clinician. More advanced studies, such as formal neuropsychological testing, EEG, neuroimaging and genetic testing, can be helpful in selected patients. In every case, the clinician must keep in mind the high concurrence rate of dementia and depression, the increasing evidence that geriatric depression may be a prodrome of dementia, as well as the research that points to shared etiologies of dementia and geriatric depression, and thus avoid taking a strict either/or approach.

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Benign Senescent Forgetfulness, Age-associated Memory Impairment, and Age-related Cognitive Decline

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Changes in cognitive functioning are prevalent in aging. These changes are perhaps best considered as falling along a continuum, with normal aging at one end of the scale and brain diseases producing bona fide dementias at the other end. It is not clear that the continuum of cognitive changes in aging and disease is entirely linear. There may be periods of plateaux in cognitive declines or even improvements in cognition as a function of adaptive brain changes. The relationship of cognitive change in normal aging and brain disease, such as Alzheimer's disease (AD) is an area of active investigation, benefiting from the advances in behavioral methods and imaging technologies^{1,2}.

Normal aging is now recognized as a complex mosaic of cognitive changes in which there is decline in some areas and a substantial degree of either stability or improvement in others³. It has been frequently observed, for example, that *fluid* abilities, those relying primarily on the efficiency of current processing (e.g. measures of spatial and reasoning abilities), exhibit pronounced and approximately linear age-related decline, whereas *crystallized* abilities, those relying more on accumulated knowledge and expertise (e.g. vocabulary measures) exhibit greater stability as a function of age. A prominent feature of normal age-related cognitive change is a decline in the speed of information processing, a decline that is more critical to the functioning of fluid abilities than to the functioning of crystallized abilities. Salthouse⁴ has developed a general theory of age-related changes in fluid cognition. The theory contains two central components: a limited time mechanism and a simultaneity mechanism. According to Salthouse, changes in these mechanisms, at an elementary level, lead to changes that are observed in a variety of cognitive tasks. For example, the time available to perform higher-order cognitive operations would be limited by an increase in the proportion of the available time occupied by initial processing stages. Similarly, if initial processing is slowed, then the products of this processing may not be available simultaneously, as required by later operations.

Between the two boundaries of normal aging and genuine cognitive impairment is a broad transition state comprising gradations of cognitive change attributable to any of a host of etiologies, including benign effects of aging to early-stage AD. Over the last three decades, a variety of nomenclatures have emerged to describe the observed transitional memory state attributed to aging. The most commonly recognized terms include "benign senescent forgetfulness", "age-associated memory impairment" and, more recently, "mild cognitive impairment".

These terms vary in subtle ways from one another. However, regardless of which terminology is used, all of these categorization schemas basically describe the same phenomenon, an admixture of mild memory problems that exceeds the range of normal cognition but falls short of being classified as dementia.

The term "benign senescent forgetfulness" (BSF) was originally coined by Kral⁵ in the early 1960s to describe a form of mild memory impairment that occurred in the context of aging and did not appear to progress to dementia. In cohorts of elderly nursing home residents followed over 4 years, Kral *et al.*⁶ observed that approximately 18% demonstrated a form of mild memory loss described as "subjective complaints of memory loss" and "difficulty in retrieving stored recent or remote information", such as names or other proper nouns. Frequently, these individuals were able to retrieve the "forgotten" information at a later time and they had well-maintained mental faculties otherwise. Compared to this group, another subgroup emerged with a more significant memory impairment, characterized by a prominent inability to retain recent information over even brief periods of time. These individuals typically had limited awareness of their difficulties and had tendencies to confabulate. This latter group, designated as "malignant memory loss", was associated with progression in symptoms to dementia, shorter survival times and increased mortality rates⁶. Kral believed that BSF was associated with physiological aging, whereas the more malignant form of memory decline was related to either vascular or degenerative disease.

The construct of BSF, although useful in succinctly describing the boundary conditions of aging, fell out of favor in later years, primarily due to a lack of standardization in the diagnostic criteria and the absence of clinical validation within representative elderly populations. The samples used in Kral's early work were criticized because of the inclusion of a very large proportion of chronic conditions, such as neurodegenerative diseases, cerebrovascular conditions and neuropsychiatric disorders. In more recent years, interest in aging and AD led to a re-emergence of attention on transitional memory states. In 1986, a workgroup convened under the auspices of the National Institute of Mental Health (NIMH) proposed the construct "age-associated memory impairment" (AAMI) to replace BSF. The new terminology had a firmer theoretical basis than its predecessor, with clinical and anatomical data to support its validity. The new nomenclature also improved upon BSF in that it included well-delineated standardized diagnostic criteria. The latter feature rendered the

new terminology more amenable to cross-laboratory investigation and potential intervention through pharmaceutical treatment trials⁷.

Briefly, the criteria for the AAMI diagnosis includes: (a) the presence of subjective memory decline in an individual over the age of 50; (b) objective evidence of memory loss on standardized testing of memory function; (c) adequate intellectual function; and (d) the absence of global cognitive decline (i.e. dementia) or other disorders that could account for the symptoms. Although similar in form to the types of criteria used in arriving at other neurological diagnoses, such as AD and vascular dementia⁸, the definition of AAMI differs in its specification of strict psychometric cut-points for the demonstration of memory loss and the absence of dementia. A patient diagnosed as AAMI must: (a) perform at least one standard deviation below the mean (for young adults) on at least one standardized test of memory function (e.g. Logical Memory from the Wechsler Memory Scale); and (b) demonstrate the absence of dementia, defined by normal performance on mental status screening using another psychometric measure, the Mini-Mental State examination (MMSE; a score of 24 or higher required).

The approach to defining the memory loss of aging based on a psychometric algorithm has been challenged on several grounds^{9,10}. One of the main issues raised is the potential for diagnostic unreliability, particularly the threat of high false-positive rates, because the criteria rely on the results of a single memory test rather than on a consistent pattern of memory deficit. Spuriously low scores on a memory test (such as on the proposed use of the Logical Memory subtest) can be obtained for any of a variety of reasons. Similarly, patients with bona fide disorders may score quite well on the test but manifest difficulties in everyday life and on more tasking neuropsychological procedures. Related to this point, the criteria do not take into account individual differences in performance (e.g. effects of low education opportunities) or any of a variety of conditions that may alter performance and result in the mistaken impression of memory deficits, such as test anxiety or mood disorders. Finally, the choice of psychometric cut-points in the criteria for AAMI are criticized as insensitive. Dementias, particularly in early but clinically diagnosable stages, cannot be satisfactorily ruled out with MMSE scores of 24 or higher. Beyond these criticisms there is an even more fundamental concern that relates to the validity of AAMI as a process distinct from AD and other dementias. There is still considerable debate as to whether serious memory loss in the absence of dementia is actually a prodromal form of AD. The provision of an acronym, along with diagnostic criteria, implies that AAMI is an independent process associated with normal aging. However, much like its predecessor, BSF, there is ambiguity in the relationship between AAMI and AD. In fact, the similarity between the two conditions with respect to the underlying pathophysiology and proposed mechanisms for drug action suggest that AAMI is either a risk factor for AD or a mild, prodromal form of the illness. Despite these limitations, the notion of AAMI has advanced studies of aging by providing uniform criteria which could be applied across multiple sites. In so doing, the AAMI construct has allowed useful cross-study comparisons of the memory loss seen in the later decades of life.

Recently, studies from the investigative team at the Mayo Institute Alzheimer's Disease Center have suggested an alternative nomenclature for patients with memory impairments of aging,

suspected to include substantial numbers of patients with preclinical AD¹¹. These investigators refer to the boundary between normal aging and dementia as "mild cognitive impairment" (MCI), and characterize its clinical characterization and outcome. Unlike the terms AAMI and BSF, MCI is not considered benign and is instead viewed as being a risk factor or transitional state between normal aging and AD. This premise has some empirical support, including incidence data suggesting that there is a significant conversion rate from MCI to AD of approximately 12–15%/year (in line with other studies suggesting 50% conversion in 5 years). Normal controls, by contrast, convert to AD on an average of 1–2%/year. Because of the care devoted towards empirically defining the borderline condition of AD and aging, the "MCI" terminology is now growing in general acceptance and is becoming the standard nomenclature used by AD investigators involved in preclinical AD studies. It is still unclear whether all patients with symptoms conforming to MCI will convert to AD. Most likely the group of patients have some heterogeneity to their symptoms, reflecting multiple physical or medical causes (e.g. cardiovascular disease, diabetes, pulmonary conditions, etc.). With continuing progress in the identification of the underlying neuronal mechanisms of cognitive decline within normal aging, more clarity will be achieved in separating the various types of age associated memory conditions. This information is critical for secondary prevention efforts in preclinical AD.

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Minor Cognitive Impairment

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Normal cognitive functioning in the elderly has commonly been conceptualized as the range of performance found in persons without identified pathology. Comparisons of mean performance between age cohorts show decrement with increasing age, which has been attributed in the past to normal physiological ageing processes. So-called ageing-related cognitive impairment, while being considered “normal”, nonetheless has been of interest to clinicians because of the physical dependency it engenders, and as such it has been given independent nosological status. A number of concepts have been proposed to describe this tail-end of the normal cognitive range, beginning with the notion of “benign senescent forgetfulness”, first proposed by Kral¹. The publication of formal diagnostic criteria for minor cognitive impairment as “age-associated memory impairment” (AAMI) was undertaken by Crook *et al.*² for the National Institute of Mental Health. AAMI refers to subjective complaints of memory loss in elderly persons verified by a decrement of at least one standard deviation on a formal memory test in comparison with means established for young adults.

As memory impairment in the elderly is more commonly observed to be accompanied by deficits in other areas of cognitive performance, an alternative concept, ageing-associated cognitive decline (AACD), has been proposed by the International Psychogeriatric Association in collaboration with the World Health Organization³. AACD refers to a wider range of cognitive functions (attention, memory, learning, thinking, language and visuospatial function), and is diagnosed by reference to norms for elderly subjects. Application of AACD and AAMI to elderly persons within the general population suggests that they are distinct clinical entities, the latter referring to a more severe state of impairment⁴.

More recently, recognition that the wide variability in cognitive functioning observed in the normal elderly may be due at least in part to the inclusion in this group of persons with prodromal dementia or other sub-clinical syndromes has led to the generation of alternative formulations, which situate minor cognitive impairment as potential pathology, for which the clinical response should be therapeutic rather than palliative. Within the 10th revision of the International Classification of Diseases (ICD-10)⁵, criteria are given for “mild cognitive disorder” (MCD), which refers, like AACD, to a broader range of cognitive disorders than memory, demonstrable by formal neuropsychological testing and hypothetically attributable to cerebral disease or damage or to systemic physical disease known to cause dysfunction. MCD is thus construed as being secondary to physical illness or impairment, excluding dementia, amnesic syndrome, concussion or post-encephalitic syndrome. MCD is also applicable to all ages, not

just the elderly. Early attempts to apply MCD criteria to population studies of elderly persons has so far met with limited success, which has cast doubt on the validity of MCD as a separate nosological entity²². On the other hand, Gutierrez *et al.*⁶, arguing for the inclusion of a similar category in DSM (“mild neurocognitive disorder”) have reviewed numerous studies implicating diverse forms of underlying pathology, in which such a nosological category would have been appropriate.

Early evidence that elderly persons with minor cognitive disorder may be at high risk of developing senile dementia has led to the development of the concept of “mild cognitive impairment” (MCI)⁷. MCI has provoked considerable interest amongst clinicians and the pharmaceutical industry, as it refers to a much larger potential therapeutic target group than senile dementia. As such, it is likely to become a more widely-adopted concept than its predecessors. The essential feature of MCI is that it is a pathological state that is potentially progressive. Beyond this, specific diagnostic criteria found in the current literature are inconsistent. Petersen *et al.*⁷ initially refer to “complaints of defective memory” and “demonstration of abnormal memory functioning for age”, with normal general cognitive functioning and conserved ability to perform activities of daily living. A later definition refers to “memory impairment beyond that expected for both age and education level”⁸. Krasuki *et al.*⁹ refer to cognitive impairment with a score of 20 or more on the MMSE, and Zaudig¹⁰ defines MCI as a score of more than 22 on MMSE or 34–47 on the SIDAM dementia scale. Others have referred to criteria based on the Clinical Dementia Rating Scale or the Global Deterioration Scale scores^{11,12}.

A central problem in the definition of MCI has been whether or not it should be confined exclusively to isolated memory impairment. Petersen *et al.*⁸ specify that in MCI general intellectual functioning should be preserved, and that only memory should be affected, as it is the restriction of the impairment to amnesic abilities which differentiates the syndrome from AD. Isolated memory impairment was observed by the authors in a series of 76 MCI subjects; however, this appears to have been part of the diagnostic criteria for entry into the study, so that the situation is somewhat circular. Apart from the general problem of cognitive domain specificity in neuropsychometric testing (i.e. ascertaining that poor performance on a memory task is purely related to memory and does not implicate other functions, such as attention and language comprehension), other researchers have noted that subjects with MCI, although primarily having memory complaints, also commonly show deficits in other cognitive domains^{11–14}. If the concept of MCI is

extended to include areas of cognitive impairment other than memory, then it may be considered to be identical to that of AACD, except for the underlying assumption that it is a potentially pathological, progressive syndrome rather than a feature of normal ageing.

To what extent may MCI be considered a prodromal phase of AD? A number of studies have suggested a significantly elevated risk of dementia in MCI subjects, with estimates of 10–15%/year of MCI subjects developing dementia to 100% over 4 years^{8,9,14–17}. These studies are, however, all small hospital-based series. Risk factors for progression to dementia derived from larger-scale studies are ApoE 4, higher age, fine motor deficit and lower pre-morbid IQ^{8,12,18}.

A number of studies have described neurological changes in CT studies of MCI that distinguish it from normal ageing and senile dementia^{9,14}. The principal characteristic observed in these studies is temporal lobe atrophy. Celsis *et al.*¹⁹ (1997) report reduced parietal-temporal perfusion and left/right parietal-temporal asymmetry using SPECT in MCI. The observed hypoperfusion levels were found to be intermediate between those found in normal and AD subjects. Julin²⁰ used aligned SPECT and MRI images to compare MCI with early AD. AD patients were found to have atrophy and cerebral blood flow (CBF) reduction in both medial temporal and temporoparietal regions, whereas MCI showed significant reduction in CBF without atrophy in the temporoparietal region only. Jelic *et al.*²¹ have used quantified EEG to demonstrate similarities between AD and MCI that differentiate both groups from the normal elderly on temporoparietal coherence and α and θ relative power. These findings suggest that MCI and AD have similar anatomical loci, with MCI being principally differentiated by the degree of impairment, and functional rather than structural change.

In a 3 year follow-up study, McKelvey *et al.*¹⁷ reported that 64% of MCI subjects had abnormal SPECT scans at baseline; 53% of this cohort developed dementia, but of those developing dementia only 67% had initially abnormal scans, giving a positive predictive value of only 50%. On the other hand, Johnson *et al.*¹⁵ have demonstrated a clear progression from MCI to dementia, based on SPECT perfusion levels from four regions; the hippocampal-amygdaloid complex, the anterior and posterior cingulate and the anterior thalamus. The authors conclude that, with semi-quantitative analysis and a spatial resolution sufficient to detect perfusion in limbic structures, it is possible to differentiate MCI from AD.

In conclusion, a number of nosological entities have been proposed to describe minor cognitive disorders occurring in elderly persons without dementia. MCI will probably evolve as one of the most important concepts in this area, with its underlying assumption that cognitive disorders in elderly persons are potential pathologies for which therapeutic care should be sought, rather than inevitable features of the normal ageing process. The concept appears to be almost identical to that of AACD, apart from its supposition of underlying pathology, but presently lacks clear operational criteria for either research or clinical application.

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Alzheimer's Disease: One or Several?

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The current research diagnostic criteria for psychiatric illness, DSM-IV¹ and ICD-10² have evolved from a clinically descriptive exercise to an operationally defined procedure with a sharp demarcation between disease categories. In studies of dementia, diagnostic criteria with a hierarchical approach to diagnosis have also been widely adopted, in which the probability of a diagnostic subtype of dementia being present in a subject is also specified³⁻⁵. Subjects who fulfil clinical diagnostic criteria for probable disease are considered to have a purer form of the disease than those fulfilling the possible diagnostic category alone. This categorical approach aims to maximize differences between cases so that they are cleanly diagnosed. Thus, the presence of a delirium excludes the diagnosis of dementia, and the presence of cerebrovascular disease debars the diagnosis of probable Alzheimer's disease (AD). So widespread is the use of these diagnostic criteria that it is now almost impossible to practise clinical psychiatry, let alone get research findings published, without reference to them.

Clearly, this approach has its uses. It enables psychiatrists from different cultures to know, within the restraints of the diagnostic criteria, what collection of phenomenology or associated pathology they are talking about. An expertise (and associated research projects) can be developed within these diagnostic categories.

However, there are problems with such an approach and these are particularly clear in the area of dementia research. Thus, a categorical approach to diagnosis can lead to the exclusion of a large and interesting group of patients who do not fall into neat divisions. Such an approach emphasizes research into differences rather than similarities, between diagnostic categories. Finally, the assumption that clinical diagnostic categories have biological meaning is questionable.

In dementing illness, mixed pathologies are common. Indeed, the presence of vascular or Lewy body pathology, in the absence of other pathology, is relatively rare⁶. The reasons why various pathologies coexist could give important clues to the aetiology of both. However, for example, examination of patients fulfilling NINCDS-ADRDA diagnostic criteria for probable AD will lead to the exclusion of patients who have vascular risk factors, and so their importance in the development of AD pathology will be underestimated⁷. Common risk factors that are likely to be important in the development of AD and vascular disease have been recently reviewed⁸, with associations found between AD and atherosclerosis, smoking, type 2 diabetes and high cholesterol. The mechanisms of such a link are not yet established. Does vascular disease play a part in the promotion of AD pathology or its presentation? Do common factors, such as insulin resistance, oxidative stress or cytokine activation, underlie both pathologies? Clearly, the examination of mixed cases in research studies would be beneficial in understanding common aetiological mechanisms.

The separation of some diagnostic categories lends an emphasis to differences where in fact greater similarities exist. Dementia and delirium, like the common subgroups of dementia, often coexist and also share many similar clinical and biological features. This finding is, ironically, emphasized by the demarcation of one

subgroup of dementia, dementia with Lewy bodies, which has clinical characteristics such as prominent attentional deficits, hallucinations and fluctuating cognition that can make differentiation from delirium, particularly in initial presentation, very difficult. Rather than emphasize differences, another approach would be to accept the clinical similarities and to examine in more detail common epidemiological factors and pathological mechanisms of action, such as acetylcholine depletion and cytokine activation⁹.

The adoption of clinical diagnostic criteria often presumes that a group of individuals, having reconciled a number of different opinions of varying political strength, have been able to define a single biological entity on the basis of its clinical features. Clearly, this is highly unlikely. Indeed, in dementia research (where we are fortunate in having generally agreed neuropathological hallmarks by which this hypothesis can be tested) it can be seen that the application of clinical research diagnostic criteria to a community-based sample of patients with dementia suggests that, while they are efficient in identifying pathology *per se*, they are notably less efficient when other pathologies are present⁶.

In summary, whilst clinical diagnostic criteria have their uses, they also have their limitations. A broader perspective is needed, which can encompass research into the common aetiological mechanisms that explain the existence of common clinical or pathological phenomena across, and between, different clinical diagnostic categories.

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Prognosis of Dementia

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Dementia occurs on a continuum with the cognitive changes of normal aging and with mild cognitive impairment (MCI)^{1,2}. Also, the process of dementia proceeds for many years after mental status assessments and conventional psychometric measures have reached floor (bottom) scores. Consequently, a clinically meaningful and useful description of the prognosis of dementia requires measures that are capable of spanning this vast cognitive continuum. Two kinds of measures that have been found to be particularly useful in describing this continuum and associated prognostic features are global assessments and assessment of the progressive functional course of brain aging and dementia. Two specific measures that have been found to be particularly useful in this regard are the Global Deterioration Scale³ and the Functional Assessment Staging (FAST) procedure⁴ (see Tables 1 and 2).

In a recent longitudinal study of more than 200 normal aged and MCI subjects followed over a mean interval of 4 years, the overall accuracy of GDS stage assignment at baseline (i.e. GDS stage 1, 2 or 3) in predicting subsequent decline to dementia was 81%⁵. In another longitudinal study of more than 100 community-residing patients with probable Alzheimer's disease (AD), who were followed over a mean interval of nearly 5 years, the correlation between change in GDS stage and time elapsed was 0.48, the correlation between change in FAST stage and time elapsed was 0.45, and the correlation between change in MMSE score and time elapsed was 0.32⁶. Consequently, the GDS and FAST measures each individually explained approximately twice the variance in AD temporal course of that accounted for by MMSE change. Furthermore, the GDS and FAST measures to some extent accounted for independent temporal variance. Consequently, these staging procedures (the GDS and FAST measures) together explained 28% of the variance (corresponding to a multiple regression correlation of 0.53), whereas change on the MMSE accounted for only 10% of variance (corresponding to a correlation of 0.32). This latter variance in the MMSE was entirely subsumed within that of the change on the GDS and FAST assessments.

What these statistics mean is that, with the staging procedures outlined in Tables 1 and 2, we can much more accurately chart the boundaries of normal aging and progressive AD and much more accurately predict the course of AD than with mental status assessments. The staging procedures also have the advantage of being clinically meaningful. This section briefly outlines the boundaries and prognosis of normal brain aging and progressive dementia, especially the dementia of AD, in terms of global, functional and traditional mental status assessments.

NORMAL BRAIN AGING

Although many, and perhaps most, persons over the age of 65 experience at least subjective cognitive complaints, many aged individuals have neither subjectively nor objectively manifest decrements in cognitive functioning. On the GDS and FAST measures, these individuals are in Stage 1. Current data indicates that the prognosis for these persons may be better than that for equivalently aged subjects with subjective complaints^{5,7}. Clearly,

the general prognosis for these persons is for continued healthy cognitive functioning.

AGE-ASSOCIATED MEMORY IMPAIRMENT

This entity is most clearly and usefully defined as one in which individuals experience subjective complaints of impairment, but in which even subtle deficits are absent upon clinical evaluation⁸. Although this condition (Stage 2 on the GDS and FAST scales), is largely benign, a recent longitudinal study found that nearly 15% of more than 100 subjects at a mean baseline age of approximately 70 years at this stage declined to a dementia diagnosis over a 4 year follow-up interval⁵. This rate of decline is higher than would be anticipated from current incidence data⁹. This rate of decline is also consistent with a recent study of Geerlings *et al.*⁷, which concluded that the presence of memory complaints in an otherwise normal elderly population is associated with an increased risk of subsequent dementia.

MILD COGNITIVE IMPAIRMENT

At the present time, the precise definition of this entity remains quite fluid¹⁰; however, the initial definition employed by Flicker *et al.*¹ remains quite useful in understanding the nature of this condition and its associated prognosis. Flicker *et al.* defined this condition as the equivalent of GDS and FAST Stage 3. This is a stage in which subtle deficits are manifest on a careful clinical interview. These subtle deficits are generally of sufficient magnitude to interfere with complex occupational and social tasks. Studies indicate that many of these subjects develop overt dementia when followed after some years. For example, Kluger *et al.*⁵ found that about two-thirds of more than 85 subjects followed at this stage manifested dementia when followed after 4 years.

DEMEMENTIA

The prognosis of dementia varies with the nature of the dementing disorder. The major form of dementia is Alzheimer's disease (AD). The prognosis of AD is described in Tables 1 and 2 in terms of progression of clinical global changes, mental status changes and functional changes. The progression of cerebrovascular dementia is generally quite similar to that shown for AD, with two major caveats: (a) cerebrovascular dementia has been shown to have a somewhat increased risk of death and morbidity than AD¹¹; and (b) overt strokes will produce a somewhat different clinical picture and course from that shown in Tables 1 and 2. Similarly, it has been noted that AD patients who develop significant cerebrovascular disease with the progression of their condition, have a significantly more rapid illness course than AD patients who do not develop cerebrovascular dementia⁶.

Lewy bodies occur in approximately 15–25% of dementia cases. However, in a great majority of these cases there is co-existing AD or cerebrovascular dementia. In these cases with Lewy bodies in addition to AD and/or cerebrovascular dementia, the clinical presentation and course appear to be those of AD or cerebro-

Table 1 Global Deterioration Scale (GDS) for age-associated memory impairment and Alzheimer's disease³ (choose the most appropriate global stage based upon cognition and function)

GDS stage	Clinical characteristics	Diagnosis and prognosis	Approximate mean MMSE ^{5,25,26}
1	<i>No subjective complaints of memory deficit.</i> No memory deficit evident on clinical interview	Normal adult	29–30
2	<i>Subjective complaints of memory deficit</i> , most frequently in following areas: (a) Forgetting where one has placed familiar objects (b) Forgetting names one formerly knew well No objective evidence of memory deficit on clinical interview No objective deficit in employment or social situations Appropriate concern with respect to symptomatology	Age-associated memory impairment (sometimes termed “normal aged forgetfulness” or “age-associated cognitive decline”). 15% develop dementia within 4 years ⁵	29–30
3	<i>Earliest clear-cut deficits.</i> Manifestations in more than one of the following areas: (a) Patient may have become lost when traveling to an unfamiliar location (b) Co-workers become aware of patient's relatively poor performance (c) Word- and/or name-finding deficits become evident to intimates (d) Patient may read a passage or book and retain relatively little material (e) Patient may demonstrate decreased facility in remembering names upon introduction to new people (f) Patient may have lost or misplaced an object of value (g) Concentration deficit may be evident on clinical testing Objective evidence of memory deficit obtained <i>only with an intensive interview</i> Decreased performance in demanding employment and social settings Denial begins to become manifest in patient Mild to moderate anxiety frequently accompanies symptoms	Mild cognitive impairment. Two-thirds develop dementia within 4 years ⁵	25–27
4	<i>Clear-cut deficit on careful clinical interview.</i> Deficit manifest in following areas: (a) decreased knowledge of current and recent events (b) may exhibit some deficits in memory of one's personal history (c) concentration deficit elicited on serial subtractions (d) decreased ability to travel, <i>handle finances</i> , etc Frequently no deficit in following areas: (a) orientation to time and place (b) recognition of familiar persons and faces (c) ability to travel to familiar locations Inability to perform complex tasks Denial is dominant defense mechanism Flattening of affect and withdrawal from challenging situations	Mild AD. Mean duration, 2 years	20
5	<i>Patient can no longer survive without some assistance.</i> <i>Patient is unable during interview to recall a major relevant aspect of his/her current life</i> , e.g.: (a) Address or telephone number of many years. (b) Names of close members of his/her family (such as grandchildren). (c) Name of the high school or college from which he/she graduated. Frequently, some disorientation to time (date, day of the week, season, etc.) or to place An educated person may have difficulty in counting back from 40 by 4s or from 20 by 2s Persons at this stage retain knowledge of many major facts regarding themselves and others They invariably know their own names and generally know their spouse's and children's names They require no assistance with toileting or eating, but may have difficulty in choosing the proper clothing to wear	Moderate AD. Mean duration, 1.5 years	14
6	<i>May occasionally forget the name of the spouse upon whom they are entirely dependent for survival.</i> Will be <i>largely unaware of all recent events and experiences in their lives</i> Retain some knowledge of their surroundings; the year, the season, etc. May have difficulty counting by 1s from 10, both backwards and sometimes forwards <i>Will require some assistance with activities of daily living:</i> (a) May become incontinent (b) Will require travel assistance but occasionally will be able to travel to familiar locations Diurnal rhythm frequently disturbed Almost always recall their own name Frequently continue to be able to distinguish familiar from unfamiliar persons in their environment Personality and emotional changes occur. These are quite variable and include: (a) Delusional behavior, e.g. patient may accuse his/her spouse of being an imposter; may talk to imaginary figures in the environment, or to his/her own reflection in the mirror (b) Obsessive symptoms, e.g. person may continually repeat simple cleaning activities (c) Anxiety symptoms, agitation, and even previously non-existent violent behavior may occur (d) Cognitive abulia, e.g. loss of willpower because an individual cannot carry a thought long enough to determine a purposeful course of action	Moderately severe AD. Mean duration, 2.5 years	5
7	<i>All verbal abilities are lost over the course of this stage.</i> Early in this stage words and phrases are spoken but speech is very circumscribed. Later there is no speech at all—only unintelligible vocalizations <i>Incontinent; requires assistance in toileting and feeding</i> <i>Basic psychomotor skills (e.g. ability to walk) are lost with the progression of this stage.</i> The brain appears to no longer be able to tell the body what to do. Generalized and cortical neurologic signs and symptoms are frequently present.	Severe AD. Mean time to demise 2–3 years; potential for survival, 7 or more years	0

Table 2 Functional assessment stages (FAST) and time course of functional loss in normal aging and AD*

FAST stage	Clinical characteristics	Clinical diagnosis	Estimated duration of FAST stage or substage in AD [†]
1	No decrement	Normal adult	
2	Subjective deficit in word finding or recalling location of objects	Age-associated memory impairment (normal aged forgetfulness)	
3	Deficits noted in demanding employment settings	Mild cognitive impairment	7 years**
4	Requires assistance in complex tasks, e.g. handling finances, planning dinner party	Mild AD	2 years
5	Requires assistance in choosing proper attire	Moderate AD	18 months
6a	Requires assistance dressing	Moderately severe AD	5 months
6b	Requires assistance in bathing properly		5 months
6c	Requires assistance with mechanics of toileting (such as flushing, wiping)		5 months
6d	Urinary incontinence		4 months
6e	Fecal incontinence		10 months
7a	Speech ability limited to about a half-dozen words	Severe AD	12 months
7b	Intelligible vocabulary limited to a single word		18 months
7c	Ambulatory ability lost		12 months
7d	Ability to sit up lost		12 months
7e	Ability to smile lost		18 months
7f	Ability to hold head up lost		12 months or longer

*Adapted from Reisberg²⁷. Copyright © 1984 by Barry Reisberg, M.D.

[†]In subjects without other complicating illnesses who survive and progress to the subsequent deterioration stage.

**Although the potentially observed duration is 7 years, patients are generally past the midpoint of this stage when brought for evaluation.

vascular dementia. Only in a small minority of dementia cases, an estimated 4% of the total, do Lewy bodies appear independently of AD and/or cerebrovascular disease. In these cases, a distinctive clinical picture has been described, marked by fluctuating cognition, recurrent well-formed visual hallucinations, and parkinsonian features¹². There is a relatively rapid progression of the dementing disorder in Lewy body dementia, in comparison with AD.

Other forms of dementia that occur earlier, as well as in later life, including frontotemporal dementia and Creutzfeldt–Jakob disease, also differ in presentation and course from the classical AD course outlined in Tables 1 and 2. This AD course, which applies to the great majority of dementia patients, is described in Tables 1 and 2 and very briefly outlined below.

Mild AD

Stage 4 on the GDS and FAST scales, this stage has a mean duration of 2 years. Although overt deficits are present on assessment, patients are still capable of independent community survival, although the ability to manage financial and similarly complex affairs becomes compromised. Patients at this stage generally endeavor to conceal their deficits, just as humans in general endeavor to appear intelligent and try to present themselves well. This concealment may also take the form of

denial, whereby the patient tries to hide his/her problems from him/herself. Another defense mechanism is a flattening of affect, in which the patient is less participatory in social situations and appears to become more quiet and withdrawn. Medications frequently prescribed for AD patients, such as the SSRI antidepressants, may mask these otherwise common symptoms of affective flattening, making these patients appear even more overtly normal, despite their cognitive deficits.

Moderate AD

Stage 5 on the GDS and FAST scales, this stage has a mean duration of 1.5 years. Patients at this stage have deficits that are sufficient to interfere with independent community survival. Patients who are left alone in the community at this stage are either assisted by neighbors, relatives or others, or they are preyed upon by less scrupulous persons in our society. Functionally, persons at this stage develop incipient deficits with basic activities of daily life. More specifically, patients begin to require assistance in choosing the proper clothing to wear for the season and/or the occasion. Without assistance, patients will, for example, wear the same clothes day after day. A variety of emotional responses develop in an attempt to cope with the deficits in this stage. These commonly include suspiciousness, anger and false beliefs¹³. The magnitude of these emotional responses is probably dependent in part on the social supports provided to the patient. Patients who perceive themselves as secure may present themselves in socially appropriate ways at this stage and may successfully conceal their deficits in social situations.

Moderately Severe AD

Stage 6 on the GDS and FAST scales, this stage has a mean duration of 2.5 years. Patients at this stage develop deficits in basic activities of daily life. First, difficulties with dressing and bathing occur. Patients will put on clothing in the improper order or backwards unless assisted. At about the same time, patients develop difficulties with independently adjusting the shower- or bath-water temperature. With the progression of this stage, problems with independent toileting and independent continence occur. Collectively, these problems are such that not only can the patient not survive independently but, additionally, spouses or other caregivers begin to require additional help to manage the patient in a community setting. Emotional problems in the patient peak in this stage and generally include aggressivity and activity disturbances¹³. The newer atypical neuroleptics can be very useful in managing the overt emotional reactions in the patient, as can psychological non-pharmacological, management approaches^{14,15}.

Severe AD

Stage 7 on the GDS and FAST scales, this is the final stage of AD. Patients succumb throughout the course of this stage; however, the mean time to demise is about 2–3 years. Patients who progress to the final substage of severe AD, may survive for 7 or more years in this stage. Speech ability breaks down prior to the advent of this stage in the course of AD, and patients emerge in the final seventh stage with very limited remaining speech; generally a half dozen or fewer intelligible words are discernible in the course of an intensive interview in which numerous queries are presented to the patient. With the progression of this stage, speech becomes even more circumscribed. Ambulatory ability may be lost prematurely; however, after speech is lost, the ability of the patient to walk is inevitably lost. Subsequently, the ability to sit

Table 3 Functional landmarks in normal human development and AD

Normal development: approximate total duration, 20 years ↑	Approximate duration in development		Acquired abilities	Lost abilities	AD stage	Approximate duration in AD (years)	Developmental age of AD	Alzheimer's degeneration: approximate total duration, 20 years ↓
	Approximate age							
	Adolescence	13–19 years	7 years	Hold a job	3, Incipient	7 years	19–13 years: adolescence	
	Late childhood	8–12 years	5 years	Handle simple finances	4, Mild	2 years	12–8 years: late childhood	
	Middle childhood	5–7 years	2.5 years	Select proper clothing	5, Moderate	1.5 years	7–5 years: middle childhood	
	Early childhood	5 years	4 years	Put on clothes unaided	6a, Moderately severe	2.5 years	5–2 years: Early childhood	
		4 years		Shower unaided	6b			
		4 years		Toilet unaided	6c			
		3–4.5 years		Control urine	6d			
		2–3 years		Control bowels	6e			
	Infancy	15 months	1.5 years	Speak 5–6 words	7a Severe	7 years	15 months–birth: infancy	
		1 year		Speak 1 word	7b			
		1 year		Walk	7c			
		6–10 months		Sit up	7d			
		2–4 months		Smile	7e			
		1–3 months		Hold up head	7f			

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up independently, to smile, and to hold up the head independently are lost. Each of the six functional substages in this final seventh stage of AD lasts a mean of a year or longer. Studies have shown continuing cognitive, neurological, and neuropathological changes over the course of this final AD stage^{16–19}. Throughout this stage, patients require continuous assistance for survival. Even with assistance, patients are very susceptible to disability. For example, physical deformities known as contractures occur in about 40% of patients in the early portion of this stage and become nearly universal as this stage progresses²⁰. Overt behavioral symptoms become less manifest in this stage and the need for psychotropic medications steadily declines¹³. Patients commonly succumb to conditions such as pneumonia, resulting from aspiration or stroke or infections from decubiti or other sources.

Retrogenesis

It has been noted that the progression of losses in many domains in AD, and also in select other dementing disorders, reverses the normal human developmental pattern, a phenomenon which has been termed “retrogenesis”^{21,22}. This process is particularly striking with regard to the progression of functional losses in AD, which precisely reverse the order of functional acquisition from birth to the adult (Table 3). Cognitive changes also reverse the normal human developmental patterns. For example, the correlation between the MMSE and mental age has been found to 0.83²³. Neurologic reflexes have been found to be approximately as robust markers of AD course as the same reflexes are useful as markers of normal infant and child development²⁴. Other physiological and pathological phenomena in AD also appear to follow a retrogenic pattern²¹. Because of these striking retrogenic relationships, the stages of AD can be usefully understood in terms of corresponding developmental ages (DAs). The management and care needs, and many AD emotional changes can be understood on the basis of the DA model²².

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Acute Management of Dementia

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Alzheimer's disease (AD) and senile dementia of the Alzheimer type (SDAT) have an insidious onset and a prolonged course (sometimes running for 20 years); so, on the face of it, the need for acute management should seldom arise.

Indeed, perhaps for many sufferers—the “silent majority”—it may not. Some old people become ever more forgetful and adapt gradually and graciously to their limitations, while their families and friends perceive them simply as starting to show their age rather than as demented. They give them a little more help every few months, which is accepted appreciatively as appropriate. The old person may receive a lot of care at home, or move to live near or with one of the family, or agree that it would be wise to go into sheltered housing or a home, until in due course a gentle death brings life to a dignified end.

However, the course in those referred to professional services—general practitioners, social workers, geriatricians and psychogeriatricians—is often less tranquil. Usually such patients have been dementing for a year or two and the referral has been precipitated by a crisis.

CRISES

Such a crisis could be when the family doctor is telephoned by anxious and irate relatives who have visited their parent over the weekend and found things worse than when they last visited 2 months previously, or the belated awareness that he/she is not coping very well and that a long weekend, like Christmas or Easter, is imminent and that there could be problems. Relatives are usually very caring and the culture supportive, but in the developed nations families are small and dispersed and both men and women are employed, while the elderly population is large—over the age of 65 in Britain¹; so to take care of a dementing elder at a distance requires considerable adaptation.

Crisis also arise where the demented people react to their disorder not with insight, but with robust denial. These are exemplars of the “Dylan Thomas syndrome”:

Do not go gentle into that good night;
Rage, rage against the dying of the light!

They age disgracefully, fighting the implications of a failing memory, mind and body every step of the way, stubbornly independent unto death unless society intervenes, either by overruling their rights or by using some form of mental health legislation. These denying demented patients are a huge challenge to the health and social services.

Traditionally, the dementias may be dichotomized into the presenile and senile forms, or Alzheimer's and non-Alzheimer's.

However, for the purposes of this chapter, the most practical division is into those who live with others and those who live all alone. The prognosis for survival of the latter, even if given good domiciliary support, is far worse².

LIVING ALONE

Demented people who live alone may do so because they are single, divorced or widowed and without children. Widowing can be an acute event, and one of the crises in dementia is when a key carer dies, leaving the dependant not only emotionally bereft but also suddenly deprived of his/her main prop. An acute grief reaction is compounded by the abrupt removal of a principal support. The work of grieving is complicated by forgetting or denying that the loss has taken place. Plans for the future may be undermined by the fitful expectation that the lost one will return.

There may be personal as well as social factors in a demented person's living alone. Some people react to the early intimations of their disorder by withdrawing and leading a very simplified, limited existence. Finding it an effort to sustain conversation with neighbours, friends and even family, they adopt an isolated, frugal life. Those who deny that they have any difficulties are unlikely to accept the help that is willingly offered. “I don't want anyone coming to my house to do my housework and shopping—poking their nose in where it's not wanted”. “Are you saying I can't look after myself? I've managed very nicely for all these years?” “Why would I want to come and live with you (or in sheltered housing, or in a home)? I've got a perfectly good home of my own, thank you very much!” These denials are often made by people with a well-preserved, assertive personality, and enough retention of language to make their wishes plain (although without the hearing, comprehension, insight or will to listen to reason!).

Living alone with so devastating a disorder as dementia is evidently risky. Accidents occur easily in those who lack the foresight or the judgement to prevent them, and demented old people are consequently over-represented in general hospital wards. Drugs needed to control diabetes, heart failure, epilepsy or arthritis may be taken erratically or not at all, with serious consequences. Malnutrition is a hazard for those who cannot remember whether they have eaten or not, to draw their money or where they have put it, where and when to shop and for what, and what to do with what they may have bought if they can find it. Cold weather adds to the dangers of falls and hypothermia. Floods and conflagrations are always possible, and failings

in personal hygiene may amount in due course to alarming squalor.

LIVING WITH OTHERS

Dementia may develop in those already living with their family—a spouse or a son or (most often) a daughter. Crises then arise from the dependant's growing infirmity and the increasing burden on the carer(s)³. Acute exacerbations of the dementia intensify the strain. These may arise from:

- The swift progression of the dementia from one stage to the next⁴. This is said to be characteristic of multi-infarct dementia⁵, but may also occur in Alzheimer's disease.
- Events affecting bodily health, such as heart failure, a urinary or respiratory tract infection, a fall leading, perhaps, to head injury; even impaction of faeces may all add to the patient's confusion.
- Depression may have a similar but more prolonged effect. Dementia is no protection against depression, especially where there is some preservation of insight⁶.
- The dynamics of the household may have altered because of some comings or goings or change in the attitudes or well-being of one of its members. Marital strain can cause, as well as arise from, disturbed behaviour in a demented member of the household.

The demented person may have moved to be with family because of increased infirmity and dependency. Occasionally, one of the family moves in with the demented person, as when a son returns home after being divorced. Such a move may be the result of some critical failure in self-care—getting lost, having an accident, being bereft, coming out of hospital unable to cope. By moving, the demented person gains safety and security, but dependency increases, autonomy dwindles, friends and familiar haunts are now at a distance and the activities necessary for daily living are much reduced. The carer has gained peace of mind at the cost of privacy and some disruption of the household⁷. The arrival of a confused grandparent, repeating him/herself, disapproving and getting in the way, may be less than welcome to the carer's adolescent children. Also, confusion in the demented is commonly aggravated by a move, so the earliest days are not the easiest.

Occasionally the strain on carers can erupt into "elder abuse"⁸. This may take the form of physical violence, as well as angry outbursts and verbal abuse. The commonest form is when there has been mutual dependency between the demented parent and the abusing son or daughter, now at the end of his/her tether and feeling trapped in the situation.

HOSPITAL, SHELTERED HOUSING AND HOMES

Demented people are, because of their accident-proneness and deficiencies in self-care, far more prevalent among the elderly in general hospital wards than in the population at large⁹. Here their problems may be exacerbated by sudden admission, hasty and inadequate communication, the discomfort and disability of whatever they have been admitted with and sometimes, unfortunately, the indifferent, dismissive, patronizing and even hostile attitudes of staff, wary of another "social admission" or "bed-blocker"¹⁰. Medication may add to confusion by lowering blood pressure, causing drowsiness, or through anticholinergic side-effects. In the setting of a busy medical or surgical ward, an apparently able-bodied but deranged older person may be perceived as a threat—disturbing sick patients by being noisy and interfering—or an undue responsibility, liable to wander off

and become lost. Consequently it is still not unknown for physical restraint—binding hands, body or feet, trapping the patient in a "geriatric" chair, using cot-sides—to be added to sedation¹¹.

A move into sheltered housing seems to have much to commend it for those who are now too forgetful and erratic to manage readily at home but not in need of full-time care. However, such a move is better made sooner than later. Otherwise, the strangeness of the new environment aggravates the confusion, and problems may arise in the use of the alarm cord or bell to call the warden, who is summoned frequently in error. Too often the stay in sheltered housing proves quite brief, before a further move into institutional care is necessary.

Although most demented people are in their own or relatives' homes, they are also major users of residential and nursing homes, some taking all comers, others specializing in the elderly mentally infirm ("EMI"). Even in ordinary homes, as many as 60–70% of the residents may be found to be demented¹². Homes which were already looking after the old person before the onset of dementia usually cope very well, but where someone is admitted because of their dementia there may be clashes because the parties have not had time to get to know each other. Demented people can, of course, be difficult, demanding and highly irrational, but sometimes "it takes two to make a quarrel" and tactless, hasty, overbearing staff may provoke escalation of a minor dispute into a major row. Other problems that may arise from communal living include fights between residents, say where one accuses the other of going off with his/her belongings or of wandering into his/her room and interfering with the bedding (which may well be true!), and antisocial behaviour such as stripping, masturbation, sexual advances, noise and disgusting eating habits. Another crisis is that the money runs out for costly care and there are urgent demands to find the resident another place!

ACUTE MANAGEMENT

Acute problems may be lessened by early identification of the dementia, taking account of how it is managed at that time, who is (or may become) the key carer and designating a key worker to guide, advise and support that person. If no key carer is identifiable among family or friends one may need to be enlisted, such as a home help or a paid "good neighbour". Such arrangements are highly dependent on good, well-organized primary health care and social services, with support from voluntary agencies and a well-resourced psychogeriatric service, all working well together. A "case manager", generally someone with a background in nursing or social services may be the best person to assemble a "package of care"¹³, but is more effective when the client is merely elderly and infirm¹⁴ than significantly demented¹⁵.

Where dementia is identified early as the result of a screening programme (e.g. for the over-75s)¹⁶, some discretion as to how, when or, indeed, whether to impart that information. There is the possibility of an adverse reaction to the label "dementia"; the family may feel that the task of caring will prove too much—beforehand they thought they were just helping someone who was ageing normally—while demented persons may be distressed by the diagnosis. However, there is the possibility of involving them with the carers in plans for their future, to prefer one kind of management to another, make a will, give an enduring power of attorney and feel that they retain some control over their own destiny¹⁷. The advent of the oral anticholinesterases for Alzheimer's disease¹⁸ demands informed consent to their use.

Key carers need respite before they feel burdened by their continuous responsibility. "Sitters-in" enable them to take a few hours out of the home alone. Meals on wheels and home helps should not be reserved solely for demented people living alone;

they ease the load on carers and help them to feel less alone with the problem. An incontinence service providing pads and collecting soiled sheets and garments for laundering is a huge help. Financial recognition of the work that has to be done in the form of an attendance allowance is often greatly appreciated, although the sum may not be great. Day hospitals and respite admissions have been demonstrated to reduce scores measuring stress and strain in carers¹⁹. Relatives' support groups, personal counselling by a community psychiatric nurse, social worker, psychologist or doctor and, where feelings in the household are running high, family therapy²⁰, may all have their place. It is important for the key worker to keep in touch with the situation, to be easily available and able to offer extra assistance, e.g. admission to hospital or a home, as and when that is needed. The credibility of the service is seriously jeopardized if the help that all recognize to be required cannot in fact be given.

An interesting field study in Cambridge²¹ randomly divided demented people living alone or at home with others into those who received routine care and those who received extra home care. Extra care made no significant difference to those living with others, but those living alone were far more likely to be in a residential home 2 years into the study if they were getting extra care than if they were not! This was less a demonstration of the ineffectiveness of community care than an indication that those who received it were correctly placed where they needed to be at the appropriate time.

The demented who live alone are especially in need of close monitoring by the key worker. Those who will accept help are obviously an easier proposition than those who will not. The home help is the chief provider of "hands-on" care, and may be required from 2 h/week to 6 h/day (at which stage, of course, the cost is not negligible). Meals on wheels not only support nutrition but also provide regular human contact, as do lunch clubs, day centres and day hospitals for those who can make their way there or be taken to them. Community ("district") nurses may get patients up, bathe them, help them to bed and give their medication—but the supply of such skilled staff is not limitless. Reliable, trustworthy volunteers are useful in befriending, doing small chores and running errands. Financial arrangements need to be made with banks, post offices, lawyers, social workers and whoever either holds the power of attorney or has been nominated as the Receiver by such a body as England's Court of Protection²².

Demented people seriously at risk who refuse help may occasionally be compelled into care or placed on a guardianship order under mental health legislation (UK)²². This, or the decision not to take such an action, is usually the result of a case conference, often convened by a social worker, and attended by relatives, neighbours, nurses, home helps and their organizers, the general practitioner, the psychiatrist and perhaps concerned volunteers, clergy and the police. Often the conclusion is that what cannot be achieved by persuasion cannot be achieved, so a close eye is to be kept on the subject of the conference until he/she becomes more willing to accept help, falls ill and goes into hospital or dies, or confounds expectations by living on in much the same state for years!

Staff in hospitals and homes need to be trained to give demented people proper care, by example and experience as well as by precept. The inculcation of respect is a good starting point—breezy, patronizing familiarity may give offence. It is important to learn not to take umbrage, to blame, to accuse of attention-seeking or provocation and to avoid futile arguments. A spacious, bright, cheerful environment and a full, appropriate programme, including some conversation, entertainment (a sing-song is much preferable to the ubiquitous television, but people should be free to opt out), exercise and bringing visitors into the regime help to prevent problems arising from boredom and inaction.

Alertness to health problems may prevent some of the exacerbation of confusion by physical illness. Depression afflicts not a few demented people, causing agitation and restlessness by day and night. A trial of antidepressant therapy is indicated, but the old antidepressants are too powerfully anticholinergic, sedative or hypotensive to be the first choice. Serotonin-specific reuptake inhibitors are to be preferred, and citalopram 20 mg daily has been shown to improve depressive symptoms in dementia in a placebo-controlled study²³. Rarely, in those who appear severely depressed, are not eating and who may have a history of severe depression, there could be a place for electroconvulsive therapy.

A full day's activities reduce sleep problems (and patients can often catch up on their sleep by day) but carers need sleep, and the use of a hypnotic for a restless patient may thus be justified. None is ideal, but among those to be considered are temazepam, a very popular benzodiazepine, 20–40 mg (which may induce a hang-over); zopiclone, a cyclopyrrolone, 3.75–7.5 mg; chlormethiazole, a short-acting drug which occasionally causes sneezing, 250–500 mg, at night; or chloral hydrate, in tablet form (the equivalent of 414 mg in a tablet), one or two at night.

There is no perfect neuroleptic in old age psychiatry: all can cause as much trouble from side effects, notably drowsiness, extrapyramidal symptoms and falls, as any benefit they bring. Demented people seem particularly susceptible to such side effects and to tardive dyskinesia²⁴. However, where the urgent control of a very disturbed patient is necessary in a setting where alternatives are not practicable, one of the major tranquilizers may be warranted²¹. The butyrophenone haloperidol, 5–10 mg i.m. or i.v. (often with procyclidine 10 mg through the same needle to prevent acute extrapyramidal reactions) is a most useful drug, and can be repeated up to 6-hourly. Once the acute crisis is over it may be replaced by one of the newer, "atypical" antipsychotics, risperidone (0.5–2 mg, twice a day), olanzapine (5 mg once or twice a day), with fewer extrapyramidal side-effects than haloperidol²⁵ and shown by meta-analysis²⁶ effectively to ameliorate symptoms of psychosis, aggression and agitation. Very popular, and reasonably safe, although quite sedative and anticholinergic, is thioridazine, 10–100 mg, up to four times a day.

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Present and Future Treatments of Alzheimer's Disease

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Modern theories of disease predict that, when valid diagnostic criteria are available and causes can be confidently ascribed to a disease condition, then effective treatments will be developed¹. In clinical practice, the diagnosis of late-onset Alzheimer's disease (AD) is not supported by definitive diagnostic findings and no consensus exists on its causes. Currently, diagnosis depends on a cluster of neuropsychological features combined with the absence of other pathologies, identified clinically or by investigation. In those patients who present at a later stage, neuropsychological deficits are so severe and global that differentiation from other aetiologies is difficult. Moreover, the presence of other causes of cognitive decline, such as stroke, does not exclude AD; indeed, the prevalence of AD is far more common in people with stroke than would be expected by chance alone.

Observational longitudinal studies suggest that both genetic and environmental factors are important in early-onset AD². Best estimates in late-onset AD are that genetic and environmental influences are approximately equal³. In early-onset AD genetic factors seem more important but at present can account for fewer than 10% of cases⁴. Known mutations that contribute to the causes of late-onset AD appear so far to be relatively infrequent^{2,5} but genetic contributions to functional ability in late life are also established⁶. Genetic susceptibility factors are less well established; apolipoprotein E polymorphisms are best known and may influence the timing of the onset of AD in almost 50% of cases⁷. For many commentators, these sparse facts encourage the search for environmental factors that certainly contribute to the onset of dementia and, by extension, to the possible modification of those exposures. These interventions may slow or even prevent the onset of dementia. So far, however, no single factor or group of factors has been reliably confirmed as an environmental contributor to the risk of dementia. This situation is similar to the problems faced by specialists in old age medicine about 30 years ago. At that time, there was some concern about the accumulation of disabilities in old age, so much so that population projections predicted increasing longevity and an increased burden posed by large numbers of disabled old people⁸. One of the great successes of modern geriatric medicine was the postponement of the expected accumulation of disability into the final year of life. This in turn has reduced the period of dependency in old age, especially for men⁹. Prevention of disability was the key outcome of the interdisciplinary research programme one "successful ageing" in the USA¹⁰. Compression of disability and morbidity has been so great that the proportion of men aged 80 years in North America and Northern Europe with at least one disability halved between 1975 and 1995¹¹.

These reductions were achieved by preventative measures to detect and reduce environmental exposures in old people known to be at increased risk for vascular disease. Hypertension, smoking, diabetes, obesity and hyperlipidaemia proved susceptible to such interventions. Likewise, smoking reduction and improvements in air quality reduced the burden posed by respiratory disease and cancer. Reduction of disease incidence and subsequent improvements in well-being in old age have prompted some researchers to speculate that much of what is regarded as "ageing" is made up of at least two processes¹². One process comprises the accumulated effects of disease and the acquired handicaps and disabilities of old age. The other is made up of at least one process, usually termed "intrinsic ageing". The nature and cause of intrinsic ageing processes are currently described using terms such as "the oxidative stress model" or "accumulated DNA damage" or "inefficient DNA repair". The formation of advanced glycated end products, the consequences of membrane lipid peroxidation and failure of immune surveillance are all included in current hypotheses concerning intrinsic ageing. However, given that the brain enjoys specific privileges, e.g. in terms of immuno-surveillance and neurons being naturally in a post-mitotic state throughout most of the lifespan, whether some general "intrinsic ageing" process applies to the central nervous system or is the same as that for other organs is questionable.

From the standpoint of dementia prevention and/or treatment, brain ageing research is now of pivotal importance to future progress. The success of modern geriatric medicine has been achieved largely by translation of epidemiological data into public health-based preventative programmes. This chapter first considers what is currently available to treat or prevent dementia. Second, it examines some of the potential for neuroprotection and dementia prevention and/or treatment provided by recent research findings in brain ageing. Third, lessons learnt from general medicine in the prevention of heart disease and cerebrovascular accidents will be considered. Fourth, future strategies proposed to slow or even prevent brain ageing and the characteristic features of AD will be briefly summarized.

CURRENT TREATMENTS

Who is Eligible for Treatment?

AD is the most frequent cause of late-onset cognitive impairment in the UK and probably affects around 350 000 people. Incidence

rates approximately double every 5 years (65–90), such that about 20–25% of those over 80 are affected. UK demographic studies predict that old people who are cognitively impaired will increase by 11% during 2001–2011. The prevalence of AD seems likely to double by 2050. All AD sufferers are currently regarded as potentially eligible for dementia treatment.

What Are the Benefits of Current Dementia Treatment?

Anti-dementia therapy benefits are now recognized to be wider than just the slowing or improvement of cognitive decline. Additional improvements include reduced disability, time to institutionalization and fewer acute medical or surgical emergencies. Disabilities place considerable burdens on health and social services and the incidence of disability increases with age. These burdens include needs for social support, especially in best use of available health services, problems of co-morbidity and institutionalization. In London, the Gospel Oak studies reported disability in 38% of community residents aged 65–74, 77% aged 75–84 and 96% aged 85+¹³ and the overall prevalence of dementia was 9.8%. Predictions based on the MRC Cognitive Function and Ageing Study (MRC CFAS) suggest that people aged 65+ with severe disability will make up 2.2% and 3.9% of the population in 2011 and 2051, respectively¹⁴. In general terms, disability from whatever cause forecasts both mortality and prolongation of length of stay after admission to hospital.

The MRC CFAS provides further valuable information on the burden of health care linked to cognitive impairment. Cognitive impairment was detected in 38% of disabled people aged 65 years and in 46% of people in institutional care. Cognitive impairment also generates other health service demands, some not immediately obvious. For example, a reduction of dementia prevalence of 1–2% would reduce the number of hip fractures in the UK by 20 000/year^{15,16}.

The Anti-dementia Drugs

Currently available drugs are based on the established cholinergic deficits in AD and early recognition that these may be sufficient to explain “core” symptoms. There are no claims that current drugs do any more than provide symptomatic relief. Large-scale randomized placebo-controlled trial results are available for three anticholinesterase drugs: donepezil^{17,18}, rivastigmine^{19,20} and galantamine^{21,22}. These show that, in general, these drugs: (a) improve global outcome; (b) slow or arrest cognitive decline; and (c) improve activities of daily living. Carers also report that some troublesome behaviours, such as apathy and apparent responses to hallucinations or delusions, are helped by drugs of this type but as yet no satisfactory clinical trial data are available. There is a consensus to support a relatively enduring good cognitive response equivalent to an arrest of disease progression of about 9 months. This is not sustained, however, and is followed by a fairly rapid decline that reverses any earlier benefit. Primarily for this reason, most commentators agree that anticholinesterase drugs do *not* modify any underlying disease process.

The generality can obscure some quite remarkable and sustained improvements in a subset of patients. Almost half of AD patients show evidence of improvement and sustain that improvement for up to about 18 months. Within this group there is a small proportion (roughly around 1 in 12) who show very extensive improvements, sometimes sufficient to permit resumption of mentally effortful recreational pursuits. Unfortunately, there is no evidence that this group of high responders obtain long-term benefits. Such patients are highly encouraging in

routine clinical practice and certainly help motivate and improve morale in dementia care teams.

This class of drugs is usually well tolerated. Common unwanted effects include cholinergic actions on the GI tract (nausea, vomiting and diarrhoea). Few data are available on the effects of these drugs on disability and institutionalization rates. Tacrine was one of the early antidementia drugs and is now discontinued. Patients receiving tacrine and remaining on doses greater than 80 mg/day may be less likely to enter a nursing home than those on lower doses or who have stopped the drug^{23,24}.

Cost Concerns

Although the licensing of cholinesterase inhibitors has introduced widely available drug treatment for AD for the first time, the costs of these drugs prohibits extensive use in less developed countries, where the greatest increase in numbers of people with AD is expected. Even within the UK there is considerable geographical variation in the availability of antidementia drugs. In part this remains attributable to poorly informed pessimism about dementia treatment, but the fact that much-needed pharmacoeconomic data were not collected in trials sponsored by the pharmaceutical industry must share part of the responsibility. Largely at the direction of regulatory authorities, the sponsors include clinically meaningful measures of competencies in daily living, and these have become the cornerstone of current recommendations to administer these drugs as widely as possible. At present, there is an impression that these drugs are “cost-beneficial” and reduce overall spending on services for dementia. If delay to institutional care is accepted as a valid proxy in economic analyses, the eventual conclusion seems likely to support their use²⁵. Data from Canada indicate that the largest proportion of costs is attributable to institutionalization. Use of donepezil for mild-to-moderate AD was associated with lower 5 year costs and less time spent with severe AD when compared with the alternative of usual care²⁶. In a retrospective cost analysis in Dutch patients with Alzheimer’s disease who were being cared for at home at the start of the study period, treatment with donepezil did not increase overall direct medical costs²⁷.

To obtain an improvement of four points on the ADAS-COG, it is estimated that between four and six patients taking donepezil 10 mg once daily for 6 months would need to be treated. Clinical experience in the UK suggests the NNT to achieve clinically significant activities of daily living score improvement is likely to be about twice that for the cognitive end point of ADAS-COG improvement of four points.

BRAIN AGEING: NEUROPROTECTION AND PREVENTION OF DEMENTIA

Neurochemical studies support an association between brain oxidative stress and AD^{28,29}. Current studies are examining whether this association is a cause or a consequence of AD, perhaps an artifact of the AD process. Therapeutic agents are currently available (and more potent compounds are under investigation) that reduce oxidative stress. These agents may prove to be potent neuroprotective agents relevant to AD.

Evidence of oxidative stress in AD is detectable throughout the brain, irrespective of the site or extent of AD neuropathology. This evidence includes increased concentrations of advanced glycation end products³⁰, nitration³¹, increased products of lipid peroxidation³² and carbonyl modified proteins^{33,34}. The sources of oxidative stress in AD extend beyond the generation of reactive oxygen species (ROS) during aerobic metabolism. Contributions are made by activated microglia near senile plaques³⁵ and by

interactions between the receptor for advanced glycation end products and β -amyloid^{36,37}. These observations have supported the hypothesis that the neurotoxic effects of β -amyloid are mediated through oxidative stress.

Therapeutic agents that reduce oxidative stress may reduce the incidence and slow AD progression. These include non-steroidal anti-inflammatory agents³⁸⁻⁴³; inhibitors of advanced glycation end product formation^{44,45} and α -tocopherol⁴⁶. The benefits in AD of *Ginkgo biloba* extract may also involve reduction of brain oxidative stress⁴⁷.

Neuroprotective actions of oestrogen in women were linked to reduced AD prevalence in some early studies⁴⁸ and prompted many subsequent observational and experimental longitudinal studies. As yet, no single study has overcome all the methodological problems associated with complex biopsychosocial questions of this type. A recent meta-analysis of 29 studies on the putative dementia-protective effects of oestrogen replacement therapy (HRT) suggested that HRT was linked with reduced risk of dementia (summary odds ratio, 0.66; confidence intervals 0.53–0.82). The authors cautioned that control for potential confounders may remove this association⁴⁹. When demographic and health confounders were taken into account, HRT was not associated with cognitive benefits in a large ($n = 1907$) US study⁵⁰ of post-menopausal women. In the UK, the general practice research database identified women born before 1950, of whom 112 481 received HRT and 108 925 did not⁵¹. Among these subjects there were 59 newly diagnosed AD cases, of whom 15 (25%) were current HRT users. Expected HRT use was estimated at 24% and the authors concluded that this type of cross-sectional observation did not support a link between HRT and protection against AD.

Raloxifene is an oestrogen receptor modulator (mixed agonist/antagonist) that is tissue selective. Potential advantages are that it does not stimulate breast or uterine tissue but is active in bone and on lipid metabolism, the last being of specific relevance to the possible involvement of cholesterol in AD (see below). Osteoporotic postmenopausal women ($n = 7478$) were entered into a 3 year multicentre randomized placebo-controlled trial of raloxifene⁵². Mean cognitive scores were similar at baseline and there were no differences between groups over the study period. The authors concluded that, in osteoporotic women, raloxifene did not modify cognitive function over time.

As yet, it is unclear whether oestrogens are involved in AD. Although early observational studies were encouraging, recent large-scale and well-conducted studies do not provide cause for continuing optimism.

LESSONS FROM THE PREVENTION OF HEART DISEASE AND STROKE

Homocysteine and Vascular Disease

There is a rare autosomal recessive condition in which very high blood concentrations of total homocysteine are associated with increased incidence of occlusive vascular disease in adolescence—even in childhood⁵³. This disease, homocysteinuria, is caused by one of several genetic defects in the enzymes that metabolize methionine; these defects occur in methylene tetrahydrofolate reductase (MTHFR) or cystathionine β -synthase (CBS). Premature vascular disease develops irrespective of the genetic defect and this indicates that homocysteine is probably responsible for the vascular damage^{54,55}. Blood concentrations of homocysteine are also increased (but to a lesser extent) in individuals who are heterozygous for either of these two enzymes, MTHFR and CBS. Inadequate intake of folic acid and vitamins B₆ and B₁₂ (co-factors in the metabolism of homocysteine) is also associated with increased blood concentrations of homocysteine⁵⁶⁻⁵⁸. Dietary

supplementation using these vitamins is an important part of the treatment of genetically determined homocysteinuria. In the general ageing population, the contributions of genetic polymorphisms and nutritional intake to the determination of plasma homocysteine concentration (and in turn the risk of vascular disease) is largely unknown⁵⁹.

Epidemiological studies indicate that there is a strong positive association between blood homocysteine concentration and the risk of vascular disease. There is now considerable interest in the possible role of increased blood homocysteine concentration in brain ageing and cognitive decline leading to AD^{60,61}. Nutritional factors are important in the maintenance of cognitive function in late life and specific dietary deficiencies may be relevant to the failure to retain mental abilities and progression to dementia⁶²⁻⁶⁶. Nutritional factors of most interest have been antioxidants, marine oils and fat-soluble vitamins. Maternal and infant nutrition is critical in neurodevelopment; folic acid is involved in the closure of the neural tube, and maternal folate deficiency during pregnancy is associated with neural tube defects. The maintenance of normal nervous system functioning in adulthood depends on an adequate consumption of B vitamins, including folate, B₆ and B₁₂. Acute deficiencies of these water-soluble vitamins are linked to neuropathy and psychosis. Age-related changes in metabolic and physiological systems may result in old people obtaining insufficient dietary folate and B₁₂ or, to an uncertain extent, through mechanisms linked to atrophic gastritis, to failure to absorb these vitamins. In turn, insufficient intake can result in the accumulation of homocysteine⁶⁰, which is associated with greater unexpected cognitive decline and poor quality of life in old people⁶⁷ and in AD⁶⁸. A recent review⁶⁰ commented on reports that low blood concentration of folate and B₁₂ is associated with poor memory, impaired and non-verbal abstract thinking in old people. Low folate concentration is associated with poor spatial copying skills and, in very old age (90–101 years), lower blood folate concentration is linked to impaired encoding and retrieval.

The mechanisms by which homocysteine could impair mental function are uncertain. Homocysteine can undergo auto-oxidation to various metabolites and ROS (free radicals) that are directly toxic to the endothelium and also negatively alter the ability of vascular muscles to relax^{69,72}. Homocysteine metabolites, such as homocysteic acid and cysteine sulphonic acid, act as endogenous agonists of NMDA receptors, over-stimulating glutamate receptors and ultimately inducing cytotoxic damage and brain cell death. Homocysteine is also associated with elevated cyclin E (a marker of cell division) in the hippocampus of AD sufferers⁶⁸. This raises the possibility that homocysteine may be involved in neuronal stimulation and trigger proliferative mechanisms, including the induction of new amyloid formation⁶⁹.

Currently, studies are under way to determine the cognitive effects in old people of strategies to reduce blood homocysteine concentrations, and early results are encouraging^{70,71}. These strategies, if effective, may prove useful in community-based interventions to prevent cognitive decline and dementia in old people.

Cholesterol Metabolism and the Risk of Dementia

There is extensive evidence to support a link between lipid metabolism and dementia. Epidemiological evidence is as follows: (a) there is an established association between apolipoprotein E allele (APOE ϵ 4) polymorphism and AD; and (b) raised systolic blood pressure and high serum cholesterol in mid-life will increase the risk of AD in late life⁷³; and (c) use of cholesterol-lowering agents (3-hydroxy-3-methylglutaryl co-enzyme A reductase inhibitors—the statins) is connected with reduced risk of dementia⁷⁴. These findings were supported by Wolozin *et al.*⁷⁵ when they observed a lower prevalence of probable AD in subjects

prescribed two different statins (lovastatin and pravastatin). Laboratory evidence also supports an association between lipid metabolism and dementia. This includes decreased processing of amyloid precursor protein when cholesterol is removed from cell culture⁶ and possible interactions between β -amyloid and low-density receptor-related protein in AD.

These observations have prompted renewed speculation along two separate lines. First, dietary manipulation of lipid metabolism may modify susceptibility to AD in subjects genetically predisposed to AD⁷⁷⁻⁸¹. Second, randomized controlled trials that compare statins known to cross the blood-brain barrier with statins that do not will prove informative in the prevention of AD. Such studies are under way at several centres.

A substantial body of epidemiological data supports a proposed causal link between blood cholesterol concentrations and risk of coronary heart disease. The relationship is somewhat unusual, in that there is a steady increase of coronary heart disease risk from the lowest to the highest cholesterol concentrations observed in free-living communities. Some rural communities in China have mean cholesterol concentrations around 3 mmol/l and mean coronary heart disease death rates about 5% of those seen in England and Wales. This observation has prompted intensive strategies to lower cholesterol in the Western world to concentrations well below those typically aimed for. Individuals not previously considered to be hypercholesterolaemic now seek cholesterol-reducing regimes in order to reduce their perceived risk of myocardial infarction. It is as yet unknown whether these greater than previously sought-for reductions in cholesterol will be associated with reduced rates of coronary heart disease, as the epidemiological data suggest. Likewise, it is uncertain whether cholesterol-lowering regimes reduce the risk of cerebrovascular accidents. Nevertheless, these general principles in the modification of serum cholesterol concentrations in health and disease seem likely to inform the use of statins in the prevention and treatment of dementia²⁸.

FUTURE STRATEGIES

The cholinergic hypothesis provided a rational basis for drug development in AD⁸³. Some symptomatic relief is consistently observed after the introduction of cholinesterase inhibitors, but subsequent attempts to improve modification of cholinergic transmission have not brought substantial improvements in rate or extent of response. There is, however, a reasonable expectation that longer-acting cholinesterase inhibitors with improved safety profiles will be available. The proposition that selective cholinergic neuronal death is a primary event in AD pathogenesis has prompted a search for therapeutic neurotrophic factors specific for cholinergic neurones.

Nerve growth factor (NGF) therapy is an example of this type of approach. It is based on extensive evidence that NGF is critical in the maturation and maintenance of cholinergic neurones and a possible modulatory role for NGF in amyloid precursor protein (APP) expression and secretion⁸⁴. So far, problems of drug delivery have largely thwarted any chance of success. NGF shares these problems with other peptide drug delivery to brain sites. In summary, these are: (a) extra CNS peptide digestion; (b) failure to cross the blood-brain barrier and achieve therapeutic concentrations in the CNS; (c) inefficient distribution within the CNS; and (d) unwanted actions in CNS. The clinical pharmacology of neurotrophic factors is in its infancy and, once mature, seems likely to offer realistic prospects of contributing to AD treatment and prevention. Uses of neurotrophic mimetics (usually small synthetic molecules with growth factor-like activity) are likely to include the promotion of neuronal survival and differentiation

after cell-based therapy⁸⁵⁻⁸⁸. Antisense strategies in AD therapy face similar problems. They possess considerable potential to target specific genes but so far none have been found useful in neurodegenerative disease⁸⁹.

Current drug development is based largely on better understanding of AD molecular pathology⁹⁰. The "amyloid cascade hypothesis" of AD postulates that amyloid deposition is critical to the development of dementia and to neuronal death. Interventions are now proposed to prevent amyloid deposition and, when it occurs, to cause amyloid to disaggregate and be scavenged from the CNS^{91,92}. Relatively little attention is paid to the development of therapies to prevent neurofibrillary tangle formation. A separate strand of research (described above) seeks to protect the brain against oxidative stress, inflammatory reactions, perturbations of calcium homeostasis, apoptosis and cerebrovascular endothelial function.

Figure 58a.1 shows key steps in the processing of APP. The deposition of extracellular amyloid is postulated to be the key event in AD pathogenesis. Three enzymes are involved; α -, β - and γ -secretases. The identification of these secretases raises the possibility that therapies aimed to modify their function could be useful in AD⁹³. Several modulators of secretase function have been described, but so far none appear to have therapeutic applications⁹⁴.

The prevention of β -amyloid deposition may prove critical in AD treatment. Senile plaques contain not only β -amyloid deposits but copper and zinc that can be solubilized by Cu/Zn chelators *in vitro*. Recently researchers in Australia and Germany⁹⁵ successfully reduced β -amyloid accumulation in APP2576 transgenic mice (predisposed to β -amyloid deposition). This raises the possibility that therapies designed to remove brain Cu/Zn may be valuable in AD.

An alternative approach to β -amyloid deposition in AD is based on the production of antibodies to β -amyloid⁹⁶⁻⁹⁹. This technique seeks to dissolve senile plaques after they have formed and has the potential to prevent their formation. In studies of transgenic mice (TgCRND8 predisposed to β -amyloid accumulation), immunization with β -amyloid antibodies produced a marked (around 50%) reduction of β -amyloid in brain and was linked to reduced cognitive impairment¹⁰⁰. More detailed behavioural studies were reported at about the same time¹⁰¹. These concluded that treated mice performed

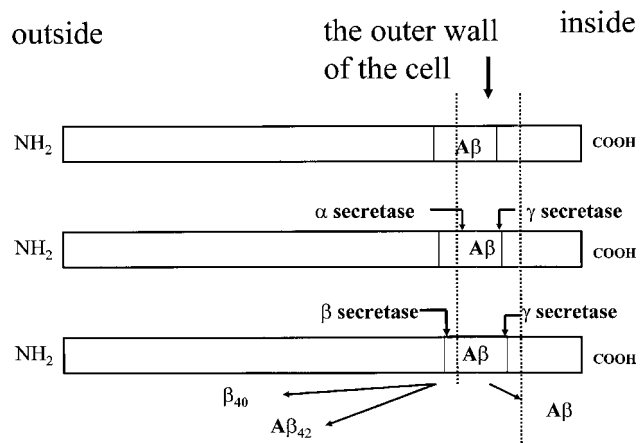


Figure 58a.1 Amyloid precursor protein (APP) is a transmembrane cellular adhesion molecule involved in synaptogenesis. It is processed by enzymatic cleavage to yield amyloid fragments (β_{50} , β_{42} and A β) of the original APP. Each cleavage enzyme acts at a specific point on APP. β - and γ -Secretases cleave at one or other end of the amyloid fragment. α -Secretase cleaves at a point between the two

“superbly” on the Morris water maze, whereas untreated mice did not. The authors remarked that their data did not support concerns that attempts to remove β -amyloid from brain might inadvertently cause wider dispersal and, therefore, more extensive β -amyloid neurotoxic damage. A later population-based observational study tested the hypothesis that the natural occurrence of antibodies to β -amyloid might reduce the incidence of dementia¹⁰². Although antibodies to β -amyloid could be detected, no reduction in AD was found.

CONCLUSION

The future for AD therapy holds much promise. As the general health of old people continues to improve and the incidence of functional disabilities is reduced, so the achievement of dementia prevention seems not only attainable but worthwhile. The huge collaborative international effort to understand the molecular pathology of AD and to design drugs to modify critical steps is certain to bring great benefit to those at risk of dementia.

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Possible Future Treatments and Preventative Strategies for Alzheimer's Disease

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PREVENTION OR CURE?

It is a truism that prevention is better than cure, although historically both are usually preceded by palliation. Whilst this seems to be true also for Alzheimer's disease (AD), in that drugs for symptomatic treatment have preceded other approaches, the situation is complicated, both because these drugs may themselves alter the disease process and because prevention and cure (in the sense of reversing or modifying the disease pathogenesis) may be inseparable. The confusion largely results from the timing of the onset of AD. At the point of diagnosis, disease has already been established, probably for many decades. Careful pathological studies have demonstrated that a marker of neuronal damage in AD—highly phosphorylated tau—accumulates in neurons in certain cortical areas in mid-life and progresses to frank neurofibrillary tangle formation and neuronal death in these areas before the onset of clinical dementia¹. Neuroimaging studies show that those with genetic risk factors for AD have evidence of functional damage decades before onset². Truly preventative strategies, therefore, preventing even the onset of AD pathology, are probably not realistic. Rather, prevention will be secondary—preventing the disease process progressing to the point where it becomes clinically manifest. This approach (prevention) differs from approaches designed to slow the progression of disease once started (treatment) only in timing. It is best perhaps to think only of disease modification; either late, in which case it looks like slowing of disease, or early, in which case it looks like prevention. From the biological point of view, both are the same.

DISEASE MODIFICATION—DIRECTIONS FROM EPIDEMIOLOGY

Identification of risk factors for AD, such as vascular factors or diabetes, points the way towards obvious potential interventions to slow disease progression. It would be hoped that reducing hypertension and head injury, improving cardiovascular health and controlling diabetes well would all reduce the conversion of early pathology to clinical dementia. Time will tell whether this is the case. Worrying for this long-term strategy is the observation that treatment with insulin carries a higher risk than diabetes alone³. This may be a surrogate for disease severity but it does highlight that the route between epidemiology and public health may not always be obstacle-free.

Other population-based studies have suggested interventional possibilities for disease modification. Most promising at present is

evidence suggesting that non-steroidal anti-inflammatory drugs may delay progression or prevent the appearance of symptoms⁴. Similarly, there is evidence linking oxidative damage to pathology and antioxidants to protection, and there may be an important role for vitamin E or some other antioxidant therapy in disease modification in the near to middle future⁵. The evidence linking hormone replacement and protection has a more limited potential, if only by virtue of gender. Studies are currently in progress that will determine whether any of these three approaches will have a clinical role in modifying the pathogenesis of AD.

DISEASE MODIFICATION—DIRECTIONS FROM MOLECULAR BIOLOGY

The advances in understanding the molecular biology of AD have been rapid and profound. It is clear already that this work will yield compounds designed to modify disease, although whether these compounds make the difficult passage between laboratory and clinic is a different matter. Preventing amyloid formation is the most obvious drug target and compounds designed to do this by inhibiting γ -secretase are in the late stages of development. Another, complementary approach is to increase the activity of α -secretase (thus hopefully reducing amyloidogenic metabolism). Interestingly, all cholinomimetic therapies should increase the activity of α -secretase indirectly, and this has been shown to be the case for M1 agonists⁶.

The other pathology of AD, tangles, is a harder target as it is not yet clear why tau aggregates. However the importance of tau aggregation as a target is emphasized by the fact that this occurs in many neurodegenerative conditions⁷, correlates with cognitive impairment and has functional consequences for neurons. Tau phosphorylation precedes aggregation and may cause aggregation in AD; certainly it inhibits tau function⁸. In neurons, tau is phosphorylated by an enzyme, glycogen synthase kinase-3 β (GSK-3 β), and inhibition of this would be expected to prevent tau phosphorylation⁹. Lithium inhibits GSK-3 β and, as predicted, reduces tau phosphorylation^{10,11}. Might lithium modify disease progression? Other GSK-3 β inhibitors are in development. GSK-3 β is also inhibited through the same signal transduction route as that induced by muscarinic agonists, and in line with this is the finding that muscarinic agonists reduce tau phosphorylation in cellular models¹². It is possible that cholinomimetic therapies (including the cholinesterase inhibitors) will have some disease-modifying effect as well as a palliative effect¹³.

Perhaps the most surprising and exciting development suggests that a vaccine is a therapeutic possibility for AD¹⁴. Amyloid peptide was used to inoculate transgenic animals overexpressing the amyloid precursor protein gene. In these animals, amyloid inoculation reduced amyloid deposition in the brain without any apparent adverse effects on the animals. It remains to be seen whether this is true in other animal models and whether cognitive deficits are also prevented. More importantly, it will be necessary to be very sure that there are no adverse consequences in man. The brain is supposedly (but probably not entirely) immunologically sacrosanct, and injecting a normal brain protein and stimulating an immune reaction has the very real potential of inducing auto-immune damage. Toxicity studies are under way that will address this concern.

TOWARDS DISEASE MODIFICATION

—A TIME SCALE

The time scale of science is not linear; progress comes quicker these days and the only reliable prediction regarding the future is that predictions will continue to be usually wrong. However, disease-modifying treatments are already in clinical trials, and strategies such as HRT, anti-inflammatories and vitamin E are in use in other contexts. Another potential disease-modifying approach is to use the cholinergic therapies that are already in use for palliation. It should be known within the first few years of the new millennium whether any of these approaches have a true benefit for patients in modifying disease and, if so, given that experience in using these compounds in other contexts is extensive, then approval for use in secondary prevention may be rapid. Other approaches modifying amyloid and tau will take longer, not least because the time taken for novel drugs to travel from laboratory to clinic is best measured in decades.

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Psychological and Psychosocial Interventions

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Psychological interventions in dementia offer attempts to ameliorate the consequences of the disorder. They do not affect the underlying pathological process but are directed at helping those afflicted to function better and more independently. Interventions of this kind are based on the implicit assumption that those with dementia remain sensitive to environmental influences, even though the extent to which this is the case may be reduced. There is now good evidence that this assumption is justified^{1,2}.

GENERAL METHODS

Psychosocial interventions in dementia have been dominated by a number of approaches, which are designed to be generally applicable to all sufferers. These carry with them, whether explicitly or implicitly, the assumption that attention to a key impairment or principle is the key to effective intervention.

The assumption that an important feature of those with dementia is that they lose their orientation to time, place and person, lies behind reality orientation (RO), which is the earliest of these general methods³. There are two facets to RO. Firstly, regular group sessions, known as “classroom” or “group RO”, involve sessions in which members may be led to recall the date and time of year, where they are, and so on. Secondly, “24-hour RO” may operate continuously and involves all who come into contact with sufferers, stressing information relating to orientation. An example of this is the nurse saying, “It is now 12 o’clock and time to take your tablets, Mrs Smith”. This emphasizes information relating to time and person and links the time to the activity to be undertaken.

Evaluative studies have largely concentrated on group rather than 24 hour RO. In a review of these⁴, it was concluded that sessions did lead to increases in verbal orientation (e.g. group members are more likely to be able to say what day it is). When it comes to changes in a wider range of cognitive functions or in behaviour, the evidence is very much less convincing.

Another general approach is that of reminiscence. This was originally proposed for normally functioning old people and is linked to the assumption that reminiscence and life review is a major task of old age. As used with those who suffer from dementia, it also builds on an area of relative strength, that of memory for the distant past. Group sessions are organized, with participants discussing how things used to be, usually with special materials such as photographs of the local city centre as it used to be 40–50 years ago. Reminiscence about how things were can be used as a basis for turning the discussion to how things are now.

Like RO, reminiscence has been popular, but relevant research has not offered strong support. The evidence that old people are

especially prone to reminisce or that it necessarily leads to beneficial effects, such as increases in well-being or mood, has proved far from overwhelming⁵. Attempts to formally evaluate effectiveness in those with dementia have produced both negative findings⁶ as well as some indications of modest benefit⁷.

A more recent development of relevance to this section is what has become known as “dementia care mapping” (DCM), which is based on the writings of Tom Kitwood^{8,9}. Kitwood stressed the “personhood” of people with dementia, and in his view the central aspect of personhood is found in social relationships with others. Good care is that which enhances personhood and well-being in terms of such things as enhancing self-esteem, enabling individuals to influence their own personal lives, promoting social confidence in terms of being at ease with others, and a sense of hope.

DCM is built more on a set of values than psychological principles, albeit values that almost no-one who is concerned for the welfare of these people would wish to dissent from, at least in general terms. What is lacking in this general approach is any detailed analysis of how these might best be turned into caring practices for elderly people with dementia, other than by feedback from an evaluative process, or “dementia care mapping” as more narrowly defined¹⁰, to assess quality of care. This is based, firstly, on coding activities or inactivities according to whether they are expressions of “well-being” or “ill-being”. Secondly, there is recording of episodes in which the person is demeaned or diminished. The obvious problems are those of reliably defining what is “well-being”, “demeaning”, etc., and no data as to the reliability of the method appear to be available.

This is just a selection of the general approaches or methods that are available and others are described elsewhere^{1,2}. Most of these suffer from the limitation that, explicitly or implicitly, they see the problem of dementia as a single issue (e.g. loss of orientation in RO) or, as in the case of DCM, offer a set of very general values and principles. The best evidence as to effectiveness is still for RO, although even the positive impact of that is limited in extent. However, the search for better methods of care is not necessarily futile, since there is evidence that different forms of intervention, such as altering the layout of the furniture to make interaction easier, or enhanced activity programmes, can produce improvement^{11,12}.

SPECIFIC METHODS

As already indicated, approaches like those described in the previous section all assume a particular key factor, whether it be lack of orientation or the minimizing of personhood, offers the

key to optimal care. Against this is the argument that even such an ultimately devastating disorder as dementia does not obliterate all individuality and, in consequence, different people with dementia will have different problems and varied needs, even if there are some common elements. This does not deny that things like DCM and RO might have a general role, but it does mean there is also a need to consider individuals and their particular problems and circumstances.

A wide range of psychological interventions directed at specific problems, such as incontinence, memory failures and social behaviour, have now been described. These are based more in the kinds of psychological treatments developed for use with other clinical populations. Space does not permit an extensive description of these and more detailed information can be gained from other sources^{1,2,13}. This section will merely offer a few illustrative examples.

Incontinence has always proved a difficult and almost intractable problem for psychological intervention. Nevertheless, some evidence of techniques able to produce beneficial effects has appeared. A system of checking for wetness and prompting going to the toilet has been found to produce positive effects, increasing the number of "dry checks"¹⁴. It is also interesting that one study has suggested that earlier failures to produce benefits with similar programmes might be at least partially attributable to staff failing to comply with the regime, rather than a failure of the method to provide positive effects when properly applied¹⁵.

Aimless wandering can also be another feature that is difficult to manage. The adaptation of principles based on operant conditioning has shown some promise as a means to reduce this behaviour^{16,17}. An interesting point in relation to this particular problem and, by implication, other problems as well, is that it is often unfortunately seen as secondary to intellectual loss and therefore only remediable if the primary problem can be tackled¹³.

Finally, one possible way of ameliorating the problem of memory loss is to use external memory aids as prompts in order to lessen the load on the individual's own memory and to support retrieval. An encouraging account has described the use of this strategy with some success in the execution of daily living tasks, such as preparing a drink or snack¹⁸.

COMMUNITY-BASED INTERVENTIONS

Most of the elderly population with dementia live in the community and are looked after by relatives, who can be under considerable strain¹. Supporting carers therefore assumes considerable importance and extensions of the approaches described above have been made to deal with problems encountered in those resident in the community¹³ (a wider discussion of community care issues is provided in Sections MI and MII).

One of the most obvious strategies is to provide support groups for carers, which may concentrate on providing information about dementia and discuss coping strategies, especially those used by group members. This strategy is described in greater detail elsewhere¹.

More formal methods, such as RO, have been adapted for use with community residents attending day hospitals. Similar effects to those obtained in applying RO in psychogeriatric wards and nursing homes were obtained. Again, the impact was most pronounced for orientation measures but with some effect on mood as well¹⁹.

Finally, it is possible to use informal caregivers, such as spouses and children, as agents to implement more specific psychological interventions of the kind outlined in the immediately preceding section^{20,21}. Problems tackled with some indication of success include the improvement of self-care skills and social interaction.

COMMENT

The most important conclusion that can be reached after surveying the evidence on psychological and psychosocial interventions for those with dementia is that people with even quite marked levels of dementia are responsive to psychological and environmental manipulation. This offers the essential foundation for the development of therapeutic or management strategies based on psychological principles.

Quite simple manoeuvres, such as rearranging chairs in groups to facilitate conversation, rather than leaving them in long regimented lines, can improve social interaction. Other relatively simple interventions can contribute to maintaining the basic skills associated with independent living. Despite being positively regarded by staff and patients, general methods like RO and reminiscence are of limited value and DCM remains to be evaluated. The best support is for RO, and indicates that small positive changes in orientation can be achieved. Since these methods are easy to apply, they can also be used as a general background on which more specific interventions can be built, and the successful addition of behavioural training to RO exploited in one investigation²² can be seen as an example of this.

It may be that an important spin-off from general methods like RO is their popularity with direct care staff. Whilst this remains to be formally demonstrated, their use may help create and maintain a more optimistic and therapeutic attitude in staff, which can be a very worthwhile achievement in itself. In turn, this should make it easier to implement more specific interventions, more focused on individual needs.

As with psychological interventions in other contexts, the best results are likely to follow from the careful functional analysis of problem behaviour, whether this be a lack of behaviour or excessive and inappropriate behaviour, with specific interventions chosen in relation to the exact nature of the problem.

Overall, psychological and psychosocial interventions in the treatment and management of those with dementia are now capable of achieving modest but useful beneficial effects. They now must be regarded as a worthwhile part of any overall management strategy.

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Informal Carers and Their Support

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This chapter is concerned with the need to support informal carers of people with dementia. It will describe how the importance of informal carers has been officially recognized in the UK, albeit partially. The shortcomings of state legislation are reflected in the narrow definitions of informal care that are still prevalent. Following a discussion of “What is caring?”, this text will describe the significance of the role played by informal carers for both individuals with dementia and at the wider level. Then the chapter will explain how the rewards of providing informal care are often overshadowed by the adverse effects on carers’ physical and emotional well-being. Some of the economic barriers faced by carers will also be outlined. Without adequate support, the ability of informal carers to maintain their role may diminish. A brief overview of the various interventions intended to “care for carers” will follow. It will be seen that many interventions are inadequate, and the need for proper evaluation will be highlighted. The chapter will end with a summary of the key messages.

POLICY BACKGROUND

In the UK the Carers (Recognition and Services) Act 1995 (1996)¹, gave some informal carers the right to ask for an assessment of their ability to care, and gave local authorities a duty to take into account the results of this when deciding what support to provide for the care recipient. However, carers can only have an assessment if they provide “regular and substantial” care and the person they care for is receiving an assessment. Thus, although the Act was a step forward, it should be extended if the true needs of dementia carers are to be met.

WHAT IS CARING?

Informal caring has been defined in many ways². On the one hand, it is often defined solely in terms of the physical activities involved, e.g.:

Anyone who looks after or cares for a handicapped person to any extent in their own home or elsewhere³.

More recently, the Carers Act (1996)¹ promoted this view with its restriction to those providing “regular and substantial” care. In contrast, some argue that whilst “caregiving” is best thought of as the physical and behavioural activity of an informal supporter, “caring” can be seen as the emotional/affective element. Nolan *et al.*² provide a very useful definition, which will set the context for the remainder of this chapter:

... caring comprises emotional, social and psychological aspects, as well as a general concern for others, in addition to practical tending.

THE IMPORTANCE OF THE INFORMAL CARER'S ROLE

The majority of caring is undertaken by relatives, and there are around 7 million informal carers of older or disabled relatives at home in the UK⁴. The extent and nature of the care provided is diverse, from continual to occasional help, and from assistance with personal care to the provision of a sympathetic ear. The number of people in England and Wales living with dementia is estimated to be around 550 000⁵. Although the number of people here caring for someone with dementia is not known, most people with dementia will have an informal carer. A study with which the author was involved⁶ found that out of 502 mentally frail older people, 68% identified an informal carer. In the USA, Haley⁷ found that 80% of the care of people with Alzheimer’s disease in 1987 was provided by relatives.

Informal caring offers enormous benefit, both to the individual (carer and care recipient) and at a wider level. Tax-payers benefit in that institutionalization of the older person is prevented or delayed where there exists a willing and able informal carer. The Carers’ National Association in the UK estimated that should just 10% of informal carers feel unable to continue their caring role, the cost of alternative care would be £2 billion/year⁸. On the individual level, many people in need of care prefer to receive it from their families, rather than formal carers⁹. However, this tendency is often dependent on the quality of the relationship¹⁰. The decision to choose formal care, on the other hand, is associated with a desire not to inconvenience or overload family and friends. Being able to provide care can be a fulfilling experience from the perspective of the carers, who often feel satisfaction at being able to improve the well-being and maintain the dignity of the older person, experience appreciation and companionship and feel they are reciprocating past help^{6,11}.

ADVERSE EFFECTS OF CARING

Despite the potential rewards of caring, many carers feel their own physical health, social and working life are adversely affected^{6,8,12}. One survey of a district health authority in England found that 14% of its workforce were providing informal care for older people¹³. Many people who are working at the same time as caring for an older relative are eager to fulfil both responsibilities. But trying to balance competing demands is often problematic, with

many carers having to work fewer hours, take unpaid leave or stop working completely¹⁴. Thus, middle-aged carers may become disadvantaged, both in the short term in terms of loss of income, and in their later years because they have relinquished pension contributions.

Moreover, psychological distress is common among informal carers^{7,15}. This is of wider concern, because psychological distress in carers is considered to be a predictor of breakdown of community care, i.e. of the older person's admission to long-term care^{16,17}. The likelihood of poor psychological well-being (depression, anxiety, psychological stress) is greater in the carers of people with dementia compared with relatives of older people without dementia^{12,18}. Vedhara *et al.*¹⁹ found that stress levels were higher in carers of people with dementia than in a control group of similar socioeconomic status.

There are inconsistencies in longitudinal research of carers of people with dementia. Several studies found no significant deterioration in psychological well-being over time in carers of people with dementia²⁰⁻²², whereas other studies found significant changes^{23,24}. It could be that depression levels are determined early in the caregiving career and remain stable for the duration of caregiving²¹. Although Wright²² found no significant changes over time in depression, carers were significantly more depressed than non-carers by two year follow-up.

TYPES OF CARE

Bowers²⁵ proposed a useful typology of caring, beginning with anticipatory care, then preventive, supervisory and instrumental care. Central to all of these is the notion of protective care, whereby carers attempt to maintain the self-esteem and autonomy of the care recipient. In many cases this clashes with instrumental aspects of caregiving. Given that formal services often provide instrumental care, conflict may occur between service providers and carers²⁶.

CARING FOR CARERS

Much of the literature has called for assessment of the needs not only of older people but also of their informal carers. The UK Carers Act (1996)¹ has answered this call, up to a point. Ideally, interventions should be "facilitative" in nature²⁶, that is, systematic, planned in conjunction with the informal carer and care-recipient, and complementing the type of care that the carer already provides. Such interventions should be evaluated using sound methodological techniques, perhaps by a partnership of academic institutions and service providers. However, not all interventions are successful in alleviating carer stress, and some carers feel that interventions create rather than relieve the pressure (the place of informal carers in the therapeutic team is covered in Chapter 123 of this volume). Various types of assistance have been designed for informal carers. These include the provision of in-home or institutional respite services, support groups, skills training and education. Although carers value such interventions, they tend to be less effective than more intensive psychosocial interventions⁷, such as counselling and psychotherapy. For a more detailed examination of a particular intervention for carers, readers may wish to refer to a special article in this book which outlines a training programme for carers of people with dementia in Sydney: see Chapter 138b.

Graham *et al.*²⁷ found that carers with greater knowledge of dementia were less likely to be depressed and more likely to perceive themselves as competent in their caregiving role. Buckwalter *et al.*²⁸ evaluated a psychoeducational nursing intervention for carers of people with dementia. In this

individual-based intervention, carers learnt how to manage behavioural problems in the care recipient, and this resulted in decreased carer depression.

However, Knight *et al.*²⁹, in a review of interventions, concluded that while the effectiveness of respite and individual psychosocial interventions is moderate, such interventions with groups of carers are weak. This can be observed in a recent study that examined the effects of a group-based, dementia carer education programme³⁰. This increased carers' knowledge of dementia, but there was no significant impact on their psychological well-being. The authors proposed that more intensive or individual interventions may be more successful. In another review of interventions, Melzer *et al.*³¹ note that the optimum type of intervention (that is, individual vs. group) may vary, depending on the needs of the individual carer. Those requiring social support may benefit more from support groups, whereas those with psychological symptoms or problems with the caring role may benefit more from individual interventions. Furthermore, a brief intervention can be of short-term benefit but may dilute over time and become ineffective. McNally *et al.*³², in their review of respite provision for carers, found little support for the existence of long-term benefits.

Zarit *et al.*³³ note that disappointing findings such as these may be due to methodological problems and/or the fact that services are provided at an inadequate level. Interventions are frequently optimistic and carers themselves not always as prepared to incorporate change as one might expect. Moreover, it is not always possible to balance the needs of carers with those of care recipients, or with cost issues, within one intervention. Thus, interventions should be facilitative in the way described by Nolan *et al.*²⁶. They should incorporate the experiences of carers, and qualitative research methods ought to be used to elicit their concerns early on in the caring career. This could prove invaluable in determining the best way to help carers, whether that be to continue caring for their relative at home or in long-term care. Social changes, such as the size and structure of the family, and increasing geographical mobility amongst the workforce already pose a threat to the availability and willingness of informal carers. Failure to extend current legislation and provide the resources necessary to ensure adequate implementation, may well result in the further depletion of this pool of carers. Finally, some economic barriers to informal caring could be removed with the provision of full-time national insurance and pension contributions for carers.

SUMMARY

This chapter has discussed the importance of supporting informal carers of people with dementia. These carers represent the foundation of physical and emotional support for people with dementia, yet the rights and needs of carers themselves are not fully recognized. Supporting a relative with dementia can be a rewarding experience but many carers suffer adverse effects on their own physical and mental well-being. Not only that, but wider social and demographic changes may jeopardize the informal caring network as it currently exists. Should these "invisible" carers become unwilling, or unable, to sustain their caring role, then the consequences would be bleak, not only for people with dementia but also in terms of the costs of alternative care. On the other hand, some would question the view that such care should be left to those, usually female, family members who may otherwise have pursued a very different way of life. Whilst moves have been made officially to recognize the importance of the carers' role, there is still a long way to go before legislation will safeguard not only their needs but also their rights, including the right not to care. But if willing carers are to remain able carers,

then adequate and appropriate support for these individuals is crucial. The problem with this is that many interventions that are meant to support carers are inadequate or have been poorly evaluated. In particular, the effectiveness of group interventions is often weak compared with individual interventions, and there has been little evaluation of long-term benefits. Innovative approaches to caring for carers are needed, but at the same time barriers to implementation, such as lack of readiness in carers to embody change or inflexibility of the socioeconomic system, need to be removed.

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The Role and Influence of the Alzheimer's Society

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Over the last 20 years there has been a quantum leap in awareness of the fact that carers of people with dementia are among the most stressed and underprivileged members of the community. This has led carers to seek each other out, support one another and form self-help groups. Self-help groups form the backbone of national Alzheimer's societies, whose aim is to use professional means to improve the quality of life for people with dementia and their carers.

It is well recognized that carers have needs which, if met, can go some way to alleviate the stress. Alzheimer's societies can be a mechanism through which carers can articulate these needs and provide and give guidance to others.

The Alzheimer's Society in the UK arose from a small beginning in 1979 to one of the fastest growing and respected voluntary organizations in the UK, with a current income of £20 million. It has helped to raise public awareness about dementia amongst all sections of the population. Its success is due to its role in defining core services, a successful fund-raising strategy and the involvement of people with personal, business and professional experience. Its particular strength is that members all belong to one national society and are encouraged to meet locally in self-help groups and other activities. This has led to a branch structure, with local committees and paid workers supported by the national body.

Dissemination of information is the most important task for an Alzheimer's society. Answering enquiries and the availability of good professional material is a proven route to greater awareness and recognition.

Carers need services. These include domiciliary care, day care and residential care. With government grants a rarity and many societies operating on a shoe-string, there are insufficient or no services in most countries. Voluntary organizations, such as Alzheimer's societies, because of their great flexibility, are in a better position to be innovative than are statutory bodies. Alzheimer's societies can provide models of good practice and promote these vigorously as a basis for government action.

Building a strong national society takes a number of years. One has to decide on the aims, look at fund-raising opportunities, make financial plans, computerize the office, get together a good set of publications, put into place parliamentary lobbying, collect a good team of staff and volunteers and decide on a branch structure. Funding research is only possible for those larger societies with significant resources. All national societies face this array of tasks. The challenge for Alzheimer's Disease International, the umbrella organization of currently 60 national societies, is to support the very varied needs of national societies in different stages of development and with different cultural attitudes to older people and voluntary help.

Psychiatrists and other mental health professionals will find it well worth their while to become members of their national society and involved in its work. This increases the likelihood that governments worldwide will take seriously the impact of dementia on the individual and family, and provide resources to support people with dementia and their families.

The Psychiatric Manifestations of CNS Malignancies

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DIAGNOSIS

A variety of signs and symptoms may lead to the diagnosis of central nervous system (CNS) cancer. One-quarter to one-third of patients will be diagnosed following a recognized seizure^{4,13,14,30}. Others will come to neurologic or neurosurgical attention because of focal weakness or headache^{21,43,44}. Tumor-related alterations in behavior will be demonstrated by 50–90% of patients at some time during their illness^{13,14,31–33}, and in as many as two-thirds of all patients, a psychiatric manifestation is the initial or only complaint^{31–39}. In a majority of these patients, the diagnosis of cancer is only made once “hard” neurologic deficits appear^{14,24,26,36,39} or at autopsy^{21,22}. Meanwhile, treatment for the psychiatric disturbance is often initiated. This pattern is most frequent in the elderly²² in cases where the tumor occurs in a relatively “silent” area of the brain (the frontal lobe, occipital lobe, ventricle, corpus callosum or septum pellucidum)^{37,40–46} and when the tumor is relatively slow-growing (astrocytoma, oligodendroglioma, meningioma)^{11,14,31,32,34,37,43,45,47,48}.

THE PSYCHIATRIC MANIFESTATIONS OF BRAIN TUMORS

Many factors, including the tumor itself, increased intracranial pressure⁶³, a variety of treatments, the patient's response to his illness and the premorbid personality, contribute to the psychiatric symptoms of patients with neoplasms (Table 60.1). These may be difficult to separate. Moreover, the signs and symptoms (both neurologic and behavioral) of a CNS cancer may be intermittent or may fluctuate^{24,41,50,51}. Behavioral disturbances may resolve with surgery or radiation therapy, even if long-standing^{39,45,48,51,52}. Frequently, they improve following ECT^{55–57} or conventional

Table 60.1 Causes of neurobehavioral disturbance in patients with cancer

Direct involvement of the nervous system with cancer focal lesions
Increased intracranial pressure
Seizures
Metabolic derangements
Nutritional deficiencies
Endocrinologic dysfunction
Opportunistic infections
Complications of therapy
Psychological response to the illness
Neurologic paraneoplastic syndromes

pharmacotherapy—tricyclic antidepressants, selective serotonin re-uptake inhibitors or methylphenidate for depression⁵⁸; lithium, haloperidol or carbamazepine for mania^{31,32,59,60}; neuroleptics for schizophrenic symptoms^{31,32,41}. Paradoxically, successful treatment of the psychiatric symptoms may delay the correct diagnosis if treatment is initiated without an appropriate search for underlying disease.

Tumor-related Symptoms

Neurologic and psychiatric disturbances related directly to the presence of cancer in the CNS can be catalogued according to the type of symptom, or the site of the lesion (Tables 60.2 and 60.3).

Behavioral symptoms are most frequently reported (75–90% of cases) with frontal or temporal lobe tumors^{33,47,61}. In about one-third of patients these are the initial manifestations

Table 60.2 Common neurologic deficits associated with focal brain lesions

Location of the lesion	Corresponding sign or symptom
Frontal lobe	Contralateral hemiparesis or hemisensory deficit; motor aphasia; Brun's ataxia; incontinence; frontal release signs (snout, grasp, etc)
Temporal lobe	Contralateral hemiparesis or hemisensory deficit; visual field abnormalities; aphasias
Parietal lobe	Contralateral neglect; visuospatial and cognitive disturbances; apraxia; contralateral visual field deficit
Occipital lobe	Contralateral visual field deficit
Basal ganglia, diencephalon, limbic structures	Contralateral hemiparesis or sensory disturbance; aphasias; visual field deficits; apraxia
Cerebellum	Limb (cerebellar hemisphere) or gait (midline) ataxia; dysarthria; eye movement disturbances
Spinal cord	Pain, weakness and sensory disturbance below the level of the lesion; sphincter dysfunction
Leptomeningeal	Cranial neuropathy; diminished (often asymmetric) deep tendon reflexes; weakness; radicular pain; sphincter dysfunction

Table 60.3 Focal central nervous system lesions associated with neuropsychiatric symptoms

Location of the lesion	Corresponding sign or symptom
Right hemisphere (especially temporal lobe and central lesions)	Delusions; schizophrenia-like behavior; mania
Temporal lobe Central lesions	Hallucinations
Parietal lobe (left) Occipital lobe (left)	
Frontal lobe Leptomeningeal Multifocal brain metastases	Delirium; encephalopathy; abulia
Temporal lobe Occipital lobe (right)	Anxiety; irritability
Frontal lobe Temporal lobe Central lesions Parietal lobe (left)	Depression
Frontal lobe Temporal lobe Central lesions Leptomeningeal Occipital lobe (left)	Memory impairment
Frontal lobe Temporal lobe	Euphoria; facetiousness

of the tumor. Distractibility, indifference, disinhibition, euphoria or mania^{43,62}, facetiousness, memory impairment^{52,53,55}, abulia, depression^{31,43,45,54,64-67,73} and confusion are common in frontal lobe tumors^{24,31,42,43,68,69}. Temporal lobe lesions can produce memory dysfunction (especially in the dominant hemisphere)⁷⁰⁻⁷², depression, and intellectual impairment^{31,61,73,74}. Anxiety attacks, irritability, dissociative states, altered sexual behavior, mania (in right-sided or bilateral lesions)^{32,60,62,75}, and schizophrenic symptoms^{24,35,46,76-79} have also been noted. The delusions which occur in this setting are usually paranoid in content and less complex in nature than those seen in true schizophrenia⁸⁰. Non-epileptic visual and auditory hallucinations develop occasionally with frontal, temporal, parietal and occipital tumors, and probably represent "release" phenomena^{82,83}. They are frequently prolonged, non-stereotyped and complex, in contrast to hallucinations produced by a seizure.

Tumors involving the limbic system, thalamus, basal ganglia, and diencephalon produce the next highest frequency of psychiatric symptoms^{33,46,84-92}. Memory deficits⁶³, including Korsakoff's syndrome⁹⁰, depression⁵³, apathy and psychomotor slowing, have been described. Delusions, visual hallucinations, disinhibition, childish behavior, mania^{59,75,91,93,94} and violent or emotional outbursts are also common⁹².

A smaller number of patients (20-30%) with tumors confined to the parietal lobe develop psychiatric symptoms^{33,61}. Intellectual impairment, depression and (with right-sided lesions) mania^{34,44,60,95} are typical. Tumors of the occipital lobe are occasionally accompanied by behavioral symptoms⁹⁶, including memory impairment, irritability and visual hallucinations^{82,96,97}. Behavioral disturbance is a relatively unusual finding in infratentorial tumors, although irritability, apathy, poor concentration and encephalopathy have all been reported^{31,46,98} and a well-characterized "cognitive affective syndrome" can occur in the setting of cerebellar tumors or tumor surgery⁹⁹.

Leptomeningeal carcinomatosis is heralded by confusion, memory loss and cognitive impairment in at least 20% of patients,

and mental symptoms ultimately develop in the majority of patients^{14,17,100-102}. While myelopathy, radiculopathy and pain are the hallmarks of spinal cord and epidural tumors, psychiatric symptoms have also been reported¹⁰³.

Seizures

Seizures are the presenting sign of primary and metastatic brain tumors in one-quarter to one-third of patients, and occur in half of such patients at some point^{14,30,104,105}. For patients with leptomeningeal disease, the corresponding frequencies are 6-7% and 14-26%^{14,16,17,30}. Tumors located in the frontal and temporal lobes are most often associated with seizures; occipital lobe foci are uncommon; lesions in the basal ganglia, brainstem and cerebellum rarely if ever produce seizures^{30,104,105}. Focal or generalized motor seizures are the most frequent and easily recognized seizure type; however, seizures may have solely behavioral manifestations¹⁰⁵. Confusion may be the only observed manifestation of a seizure arising from any location. Visual and olfactory hallucinations may arise from frontal, temporal or occipital lobe foci. Memory lapses¹⁰⁶, feelings of anxiety or fearfulness¹⁰⁷, aggressive, inappropriate or psychotic behavior¹⁰⁸ and distortions of sound, space or size occur with temporal and frontal lobe seizures¹⁰⁹⁻¹¹³. Rarely, the sensation of fear can be so overpowering that patients will run from a vaguely perceived threat ("cursive" epilepsy)¹¹⁴. Visual, auditory and olfactory hallucinations may be poorly formed (flashes of light, hissing or buzzing, unpleasant smells) or quite elaborate¹¹⁵. Patients may describe familiar (*déjà vu*) or unfamiliar (*jamais vu*) pictures or situations, snatches of music or overpowering (but inappropriate) feelings¹¹². Disinhibition and feelings of compulsion¹¹⁶ can occur with frontal lobe seizures. Schizophrenic^{101,116} and manic-depressive symptoms¹¹²⁻¹¹⁷ have been reported with temporal lobe foci in the dominant and non-dominant, hemispheres, respectively. Depression, weeping or laughter ("gelastic" epilepsy) also occur¹¹⁴⁻¹¹⁸. Although controversial, interictal behavioral abnormalities probably develop in a higher percentage of epileptic than non-epileptic patients¹¹⁹⁻¹²⁴.

Paraneoplastic Disorders

Psychiatric disturbances can also be produced by a number of indirect effects of cancer on the nervous system (Table 60.1). Of these, the neurologic paraneoplastic syndromes are the most difficult to diagnose¹²⁵⁻¹²⁹. These are seen much more frequently in the setting of systemic cancer than with primary brain tumors, and produce identifiable syndromes resulting in profound neurologic disability, in the absence of other causes. One type of "remote effect", paraneoplastic encephalomyelitis (PEM), frequently produces behavioral symptoms¹²⁹⁻¹³². PEM is an inflammatory disorder of grey matter that may involve any level of the CNS, including the limbic system, cerebellum and spinal cord. With limbic encephalitis, the gradual (average 10.5 months) onset of anxiety, depression, hallucinations, bizarre behavior, paranoia and marked impairment of recent memory, progressing to dementia (the "Ophelia syndrome")¹³², are characteristic. In one-third of cases, behavioral symptoms precede the diagnosis of cancer. Small cell carcinoma of the lung is the most common underlying malignancy, followed by breast, ovarian, gastric, testicular, uterine and non-small cell lung cancers and Hodgkin's disease. In addition to a characteristic clinical presentation and setting, well-defined serum antibody markers (Hu, Ma/Ta, CV2) are often present¹³³⁻¹³⁵ and characteristic MRI findings have also been described^{136,137}.

Effects of Treatment

A final very important contributor to the psychiatric morbidity of patients with brain tumors is the effect of treatment¹³⁸. Because patients are living longer and treatments are becoming more aggressive, the psychiatric complications of treatment have become common. While these "late effects" have traditionally been blamed on cranial irradiation, concurrent chemotherapy, corticosteroids and surgery are important contributors. The best-studied late effect is cerebral radionecrosis¹³⁹. Typically, an enhancing mass develops at the site of previous cranial irradiation several months to several years after the completion of treatment. While headache, seizures and focal neurologic defects are often present, insidiously progressive personality change, abulia and lethargy may be the only early manifestations. A second late complication of cranial irradiation, also developing months to years after the completion of treatment, has been termed "radiation-related dementia"⁴⁰. This is a more diffuse brain process, betokened radiographically by cortical atrophy, ventricular enlargement and increased white matter signal on T2-weighted and FLAIR MRI images. Again, gradually progressive cognitive impairment, abulia, short-term memory loss and personality change occur. Focal neurologic deficits are uncommon, although gait impairment and incontinence may develop. Frequently the correct diagnosis of a treatment-related complication is delayed while the diagnoses of depression, Alzheimer's disease, Parkinson's disease or normal pressure hydrocephalus are considered.

APPROACH TO THE PATIENT

The need for integration of psychiatric, neurologic and oncologic insights arises in at least three diagnostically different settings in patients with CNS cancer. In the most common scenario, a patient with known cancer or a proven primary brain tumor develops new or progressive psychiatric deficits. Frequently the symptom is iatrogenic, and withdrawal of the offending agent or substitution of some other therapy will be of benefit. Sometimes recurrent or progressive disease (a new metastasis, regrowth of a treated brain tumor) is the cause, and antineoplastic therapy is indicated. The new onset of seizures or a paraneoplastic disorder may be at fault, the symptom may have evolved from the patient's reaction to his illness. Recognition of these possibilities and an appropriately directed evaluation will often be of diagnostic and therapeutic value and will improve both the length and quality of life.

The patient with known psychiatric disease who develops new symptoms represents a second type of challenge. If focal neurologic deficits are prominent, a vigorous evaluation usually ensues. If the earliest or most obvious signs and symptoms are behavioral, a malignant etiology may be overlooked. Because psychiatric disease and cancer are both common, the chance development of cancer in a behaviorally abnormal patient will account for some of the apparent excess of brain tumors arising in patients in psychiatric hospitals. However, because behavioral changes are a common manifestation of CNS tumors and are especially easy to overlook in patients with chronic psychiatric illness, many cases elude early diagnosis. In patients with longstanding psychiatric illness, therefore, periodic evaluations should also include a detailed neurologic assessment.

A third diagnostically difficult situation occurs when an elderly patient presents with new psychiatric symptoms and no known cancer. Submitting all such patients to periodic neuroradiographic, electroencephalographic, neurologic and laboratory evaluations would eliminate most misdiagnoses, but is impractical and costly. Most secondary psychiatric disturbances in the elderly are due to toxic-metabolic, endocrine, cerebrovascular or

infectious etiologies⁴¹. A detailed history supplemented by a few simple laboratory tests is usually adequate for diagnosis of these disorders. In most patients with CNS cancer, at least subtle neurologic abnormalities are demonstrable at the time of psychiatric presentation, although a careful neurologic examination may be required^{47,52}. If not attributable to another known etiology (e.g. stroke, trauma, multiple sclerosis), such abnormalities should prompt additional studies. The presence of seizures, evidence of increased intracranial pressure (papilledema, headaches, nausea and vomiting), a disturbed level of consciousness, gradual intellectual decline, "frontal lobe" findings (also seen with temporal lobe and deep cortical lesions; Table 60.2), or persistent, unexplained headaches should also trigger further evaluation. In such patients, neuroimaging or examination of the cerebrospinal fluid (if leptomeningeal disease is suspected) is the best diagnostic approach.

Elderly patients with the new onset of psychoses, mania or hallucinations, suggestive family histories, normal neurologic examinations and no neurologically worrisome complaints rarely harbor CNS malignancies. Nevertheless, the number of these "idiopathic" cases, after toxic and metabolic etiologies have been excluded, will be so few and so unusual that neuroimaging is probably justified. In contrast, depression is common in the elderly. A careful history and examination are obligatory.

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Peripheral Neuropathy and Peripheral Nerve Lesions

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Peripheral neuropathy is a diffuse process involving either sensory nerves, motor nerves or, more frequently, both. It is the most common neurologic disease of the elderly. The known causes of polyneuropathy are numerous and are frequently associated with other systemic illness (Table 61.1). Many occur in patients with psychiatric symptoms. Additionally, drugs must always be considered as possible etiologies.

In addition to diffuse peripheral neuropathy, the pattern of single peripheral nerve involvement is termed mononeuropathy. The causes include acute trauma or entrapment but often are non-traumatic in origin, presumably representing either chronic trauma or due to other factors, such as infarction of the nerve (diabetes mellitus or collagen vascular disease). Patients with many psychiatric diseases are more prone to these problems, i.e. peroneal or ulnar pressure palsies in depressed patients.

POLYNEUROPATHY

Symptoms of diffuse peripheral neuropathy include a “stocking glove” distribution of sensory loss usually involving the feet and later the hands. With sensory involvement, patients complain of numbness and tingling or pain in the feet. Early on, the hands are rarely involved to any significant degree. On examination, varying degrees of decreased perception to pain, temperature or vibratory sense and, less often, joint position sense may be seen in the distal extremities, particularly in the feet. When this sensory loss is severe in the lower extremities, there may be unsteadiness of gait, worse with the eyes closed.

When the motor nerves are involved, distal muscles of the feet and hands may be weak. The patient may walk with a “foot drop” or be unable to stand on tiptoe due to distal lower extremity weakness. Intrinsic hand muscle weakness produces decreased hand grip. Reflexes are hypoactive, initially at the ankles, but may subsequently be absent at the knees and even in the upper extremities. All symptoms and signs, including the reflexes, are usually symmetrical.

The temporal profile of the symptoms may suggest possible etiologies. Neuropathy of sudden onset is seen in inflammatory or vasculitic disease, whereas patients with familial history and long-standing neuropathy are more likely to have hereditary disease.

We will discuss only the most common diffuse neuropathies.

Landry–Guillain–Barré–Strohl Syndrome (AIDP)

Immune polyradiculoneuropathy (polyradiculoneuropathy, Guillain–Barré syndrome) may be preceded by a minor febrile illness or following immunization. Symmetrical motor weakness, often beginning in the legs and ascending, and areflexia with minimal sensory involvement, are the primary features. The cerebrospinal fluid (CSF) protein is usually increased and no cells are present. This relationship is termed cytoalbuminologic dissociation. Facial diplegia without extra-ocular muscle or pupillary involvement may occur. Respiratory weakness may develop rapidly. The severity of the illness ranges from minimal weakness to flaccid quadriplegia. Although usually idiopathic, known causes include AIDS, rabies, cytomegalic and other viruses. Porphyria may cause neuropathy along with psychiatric symptoms.

Quadriplegia may develop rapidly, requiring respiratory support within an hour. Usually the symptoms progress over 3–4 days, although occasionally it may take weeks to develop the maximum deficit. Spontaneous recovery, usually complete, occurs over weeks or months as a rule. However, some patients have serious residual deficits.

Respiratory function may deteriorate rapidly, even without obvious respiratory distress. When Guillain–Barré syndrome is suspected, serial pulmonary functions should be performed. When the forced vital capacity drops below 1–1.5L, intubation is indicated. Autonomic dysfunction is common, with symptoms ranging from bladder dysfunction and labile blood pressures to cardiac rhythm disturbances.

Therapy is supportive. Maintenance of adequate respiration and nutrition, treatment of infection and autonomic disturbances, good nursing care and physical therapy to prevent contractures are required. Plasmapheresis, immunoglobulin therapy (IgG) and, less often, steroid therapy may shorten the course of the illness.

A chronic relapsing form of inflammatory demyelinating neuropathy (CIDP) is an increasingly important neuropathy and often responds to corticosteroid treatment. Nerve biopsy is occasionally helpful to confirm the diagnosis.

Diabetic Neuropathy

Various portions of the peripheral nervous system can be affected by diabetes mellitus. Diabetes is common in the elderly. Individuals with psychiatric disease will frequently have diabetes with neuropathy concomitantly. The most frequent pattern is a distal symmetric predominantly sensory polyneuropathy. Painful

Table 61.1 Peripheral neuropathy: etiology and classification

1. Neuropathy associated with toxic metabolic states
 - A. Vitamin deficiency (B₁, B₁₂, B₆, nicotinic acid, pantothenic acid and folic acid)
 - B. Diabetic
 - C. Uremic
 - D. Hepatic
 - E. Thyroid disease
 - F. Dysproteinemias and paraproteinemias
 - G. Alcohol
 - H. Amyloid
2. Inherited peripheral neuropathy
 - A. Charcot–Marie–Tooth
 - B. Dejerine–Sottas (hypertrophic)
 - C. Roussy–Lévy
 - D. Refsum’s disease
 - E. Tangier disease and A β -lipoproteinemia (Bassen–Kornzweig’s)
 - F. Neuropathy associated with the leukodystrophies (metachromatic leukodystrophy, Krabbe’s disease, adrenoleukodystrophy)
 - G. Fabry’s disease
 - H. Porphyric neuropathy
 - I. Friedreich’s disease
 - J. Tomaculous neuropathy
3. Infectious, inflammatory and post-infectious neuropathy
 - A. Guillain–Barré syndrome, inflammatory polyradiculoneuropathy
 - B. Diphtheria
 - C. Leprosy
 - D. Sarcoid
 - E. CIDP (chronic inflammatory)
 - F. AIDS
 - G. *Campylobacter*
4. Neuropathies associated with malignancy
 - A. Neuropathy associated with lymphoma and Hodgkin’s disease (sensory, motor, mixed)
5. Toxic neuropathies
 - A. Heavy metals
 1. Lead
 2. Arsenic
 3. Mercury
 4. Thallium
 - B. Toxins
 1. Acrylamide
 2. Trichloroethylene
 3. Benzene
 4. Carbon tetrachloride
 5. TOCP (triorthocresyl phosphate)
 - C. Drugs
 1. Vincristine, vinblastine
 2. Chloroquine
 3. Nitrofurantoin
 4. Phenytoin
 5. Disulfiram
 6. Isoniazid
 7. Thalidomide
 8. Excessive B₆ administration
 9. Dapsone
 10. Amioderone
 11. *cis*-platin

burning or numbness and tingling in the toes and feet, and less commonly the hands, are present. Often a “stocking glove” distribution of decreased pain, temperature and vibration perception is present, along with decreased distal reflexes.

Patients with longstanding diabetes or poor glucose control usually have more severe neuropathy, although these do not always correlate well. In addition, mononeuropathy (involvement of single motor or sensory nerves) is frequently seen in diabetes.

Femoral neuropathy produces pain in the anterior medial thigh, with weakness in the proximal muscles of the leg. Lumbosacral plexus involvement with weakness and atrophy of the thigh muscles is termed diabetic amyotrophy. Mononeuritis multiplex involves multiple sensory and motor nerves in an asymmetrical fashion, usually due to infarction of the nerves. However, multiple mononeuropathies may also occur, due to pressure or entrapment. These need recognition so that patients can be taught how to prevent further trauma or compression of their nerve. Radiculopathy in the thoracic or lumbar area without disc herniation also occurs frequently in diabetes. In the thoracic area, patients complain of chest wall or abdominal pain, usually unilaterally, and may have abdominal musculature weakness. The autonomic nervous system may also be involved, causing orthostatic hypotension, bladder and gastrointestinal disturbances, skin changes, sweating abnormalities and impotence. A penile prosthesis may be helpful in male patients with impotence.

Alcoholic Neuropathy (Nutritional/Toxic)

This neuropathy is most often sensory, with pain on the soles of the feet and loss of ankle reflexes. Minor motor involvement may occur, but is seldom severe. This is associated with vitamin B₁ deficiency, but is also likely a direct toxic effect from chronic alcohol use. A history of the amount and frequency of alcohol intake is essential in evaluating these problems. Abstinence from alcohol is necessary to result in any improvement. Psychiatric care is usually the predominant need in these patients, however.

Evaluation of Patients with Neuropathy

Evaluation for systemic disease in patients with peripheral neuropathy must include several factors: patient history, physical examination, basic laboratory studies and special laboratory analysis.

The history should include a detailed family history, but this may be difficult to obtain. Sometimes additional family members must also be examined. The social history should include occupation, types of hobbies and possible exposure to toxins. Other medical illness should be noted. A complete record of all drugs used, including prescription, non-prescription and recreational drugs, should be obtained. One should determine whether onset of symptoms correlates with initiation of a drug regimen. Recent medical history should include risk factors for possible infectious etiologies, such as tick exposure, high-risk sexual

Table 61.2 Approach to history in peripheral neuropathy

- Family history:* diabetes mellitus, pernicious anemia, amyloidosis, porphyria, Refsum’s disease, Tangier disease
- Social history:* alcoholism, occupation or hobbies with possible toxic exposure (carbon tetrachloride, carbon disulfide, carbon monoxide, trichloroethylene, trinitrotoluene, benzene, *o*-dinitrophenol, lead, arsenic, bismuth, mercury, thallium, copper, silver, gold, antimony, zinc), intentional poisoning (suicide, homicide), heavy smoking (carcinoma of lung)
- Medication history:* sulphonamides, emetine, hydralazine, nitrofurantoin, diphenylhydantoin, glutethimide, isoniazid, allopurinol, thalidomide, insulin
- Recent medical history:* infections (AIDS, diphtheria, tuberculosis, infectious mononucleosis, infectious hepatitis, syphilis, typhoid, typhus, “strep throat”, cat-scratch fever), malignancy (direct invasion or remote effects), gastrointestinal disturbances (seen with arsenic, lead, porphyria, thallium, vitamin deficiencies, pernicious anemia, hepatitis, Tangier disease)

Table 61.3 Physical examination in peripheral neuropathy

Hair: alopecia (thallium, arsenic), premature graying (pernicious anemia)
Skin: dry skin (myxedema), dermatitis of exposed surfaces (pellagra, porphyria), erythematous sweaty palms (alcohol, arsenic), depigmented areas (leprosy), ecchymoses (blood dyscrasias, hypercortisonism), butterfly rash or poikiloderma (lupus), ichthyosis (Refsum's disease)
Nails: Aldrich–Mees lines (arsenic), petechiae under nails (subacute bacterial endocarditis)
Gums: lead line (lead), gingival hyperplasia (diphenylhydantoin)
Tongue: glossitis (pernicious anemia, pellagra), large tongue (acromegaly, amyloid)
Sore throat: infectious mononucleosis, diphtheria, leukemia
Salivary glands: enlargement (sarcoid)
Eyes: third-nerve palsy without pupillary involvement (diabetes)
Fundi: Roth's spots (subacute bacterial endocarditis, blood dyscrasia), hypertensive changes (uremia, periarteritis), dilated veins (macroglobulinemia), retinitis pigmentosa (Refsum's disease)
Adenopathy: infection, malignancy, sarcoid
Cardiac enlargement: myxedema, beri beri
Hypertension: periarteritis, porphyria, thallium
New heart murmurs: subacute bacterial endocarditis
Hepatomegaly: alcoholism, malignancy, hepatitis
Ankle edema: uremia, beri beri, malignancy of kidney (especially if unilateral)
Wristdrop: lead
Slow relaxation of deep tendon reflexes: myxedema
Joint deformities: rheumatoid arthritis with or without vasculitis, gout
Reddish brown urine: porphyria

Table 61.4 Initial laboratory studies in evaluation of peripheral neuropathy

Urinalysis: chronic nephritis, hematuria (malignancy, proteinuria, myeloma)
Erythrocyte blood count and hemoglobin: anemia (infection, malignancy), macrocytic anemia (pernicious anemia, diphenylhydantoin), basophilic stippling (lead), acanthocytosis on fresh smear
Blood smear, fresh: acanthocytosis
Leukocytes: depressed count (lupus, toxicity), increased count (periarteritis, infection), hypersegmented neutrophils (pernicious anemia), abnormal leukocytes (leukemia, malignancy, toxicity, infectious mononucleosis)
Sedimentation rate: elevated in collagen disease, infection, malignancy, cirrhosis
Chest X-ray: carcinoma of lung, sarcoidosis infection, enlarged heart (myxedema, beri beri, hypertension)
Serum creatinine: uremia (including lupus)
Fasting blood sugar (glucose tolerance test): diabetes mellitus
Thyroid panel: myxedema
Serum cholesterol: decreased in thyrotoxicosis and Tangier disease, increased in myxedema

Table 61.5. Additional laboratory studies (if clinically indicated)

Genetic studies: (GM1, H5MN)
Antibodies: (anti-maG, anti-Hu)
HTLV I, HIV (serum)
Hair and nail analysis: arsenic
Urine: heavy metals, porphyrins
Liver function studies: hepatitis, alcoholism
Schilling's test: pernicious anemia
Cultures: infection
Serum uric acid: gout
Rheumatoid factor
Lupus erythematosus preparation (FANA)
Heterophil antibody titer
Biopsy of nodes, liver, kidney, testicle (periarteritis): malignancy, sarcoid, lupus, cirrhosis
Rectal biopsy: amyloid, Tangier disease
Nerve biopsy: vasculitis, amyloidosis, leprosy, sarcoidosis, embolic disease, Refsum's disease

Table 61.6 Genetic (DNA) tests in peripheral neuropathy

- Hereditary
 CMT1 Evaluation (DNA) CMT 1A, 1B, 1X
 Peripheral myelin protein 22 (PMP22)
 Early growth response 2 gene (EGR2)
 Axonal HMSN, Dejerine–Sottas, congenital hypomyelination, HNPP, Refsum's
 TTR 30 Amyloidosis DNA Test
- Acquired peripheral neuropathy:
 MAG, GM1 Triad, Sulfatide, Galop,
 Hu, MAG "Dual" Antigen
 Autoantibodies for GM1, MAG, GQ16,
 Sulfatide, Galop

activity or blood product exposure, and include a detailed review of systems (Table 61.2).

In addition to the neurologic examination, one should include attention to the general examination for evidence of systemic illness that could cause the underlying neuropathy (Table 61.3). Initial studies in the evaluation of peripheral neuropathy include many routine laboratory tests (Table 61.4). Specific laboratory studies are also useful. Nerve conduction studies reveal diffuse abnormalities in the motor and sensory nerves in patients with diffuse neuropathy. Electromyography is abnormal if axonal damage has occurred. These tests characterize the neuropathy as primarily demyelinating or axonal and recognize unusual patterns of response. Specific patterns are useful in establishing an etiology².

Genetic studies, analysis of hair, nails, and urine for heavy metals, liver function studies, Schilling's test, cultures, fluorescent antinuclear antibody (FANA), sedimentation rate, rheumatoid factor, evaluation for malignancy and possibly nerve biopsy, may be required in specific cases (Tables 61.5 and 61.6).

COMMON MONONEUROPATHIES

The most common mononeuropathies include involvement of the median, ulnar and radial nerves in the upper extremity and femoral, peroneal, lateral femoral cutaneous and sciatic nerves in the lower extremity. In addition, truncal neuropathies and even cranial nerve involvement may cause a localized sensory or motor deficit (Table 61.7).

Table 61.7. Common mononeuropathies

Cranial mononeuropathy
 VII Facial (Bell's palsy)
 V Trigeminal (mental nerve)
 IV, VI, II, III

Extremity mononeuropathies
 Median
 Ulnar
 Radial
 Brachial plexus
 Lateral femoral cutaneous (meralgia paresthetica)

Femoral
Peroneal
Obturator
Sciatic

Truncal neuropathy
 Intercostal (*Herpes zoster* and diabetic)
 Posterior primary rami T2–T6 (notalgia paresthetica)

Knowledge of the symptoms, signs and anatomic distribution of specific peripheral nerves is essential to recognize and diagnose the mononeuropathies. These mononeuropathies are common in the elderly. The most common chronic mononeuropathy is carpal tunnel syndrome (CTS), entrapment of the median nerve as it passes under the carpal ligament at the wrist.

Entrapment injury to the ulnar nerve is common in chronic illness, particularly in the elderly population. Psychiatric patients restricted to bed or a wheel chair use their elbows for support and often compress the ulnar nerve at the olecranon notch, producing a "tardy ulnar palsy". The supine position, with the arms in a position of mild flexion with the forearm pronated, exposes the ulnar nerve to chronic pressure.

The radial nerve innervates the extensor muscles for the fingers, wrist and elbow. The most common site of injury of the radial nerve is in the proximal portion of the nerve as it wraps around the humerus. Injury often occurs when the patient is deeply asleep or unconscious (usually due to alcohol or sedation) while the arm is held in abduction and lateral rotation. Injury results when the nerve is compressed by the head of a sleeping partner lying against the humerus ("honeymoon palsy"); or falling asleep with the arm propped over a bench, chair or bar ("Saturday night palsy").

The lateral femoral cutaneous nerve, a sensory nerve that arises from the lumbar plexus, is often entrapped at the medial border of the anterior superior iliac crest, producing an acute or subacute onset of numbness and a disagreeable prickly sensation over the lateral thigh (meralgia paresthetica). Sigmund Freud was one of

three early describers of this and experienced it himself. Recognition is important, if only to avoid intervention to treat other conditions, such as lumbar radiculopathy.

Peroneal nerve injury produces footdrop or loss of sensation in the anterolateral surface of the foot. This can be subtle. The most common cause of common peroneal nerve injury is crossing the legs. Immobility is a predisposing cause, and depression, stroke or dementia is present among many patients.

Sciatica is usually caused by a protruded lumbar disc affecting the S1 root. However, the sciatic nerve is susceptible to injury at various sites in the buttocks causing identical symptoms and signs.

Idiopathic inflammation of the brachial plexus (neuralgic amyotrophy, cryptogenic brachial plexopathy, Parsonage–Turner) predominantly affects the superior trunk of the brachial plexus. A deep aching pain in the axilla or shoulder is followed within a few days by weakness of muscles supplied by the superior brachial trunk.

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Electroencephalography (EEG)

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It is 60 years since Berger¹ demonstrated electroencephalographic abnormalities in an histologically confirmed case of Alzheimer's disease. Since then, the electroencephalograph (EEG) has been widely used in the investigation of patients with suspected organic brain disease. To interpret the results in an individual patient, a number of observations relating to the nature and origin of the EEG signal and the impact of healthy ageing on its waveforms must be considered.

PRINCIPLES OF EEG INTERPRETATION

The EEG samples the electrical activity of the cerebral cortex only. Further, widespread synchronous involvement (6cm² or more) of the cortex is required to significantly alter the scalp EEG waveforms. Cerebral neuronal dysfunction caused by different disease processes produces essentially similar patterns of EEG disturbance. Hence, with a few exceptions, the type of EEG change cannot predict the precise nature of the underlying pathology. Finally, acute cerebral disease or rapidly deteriorating dysfunction cause the most florid electrical disturbance, while static lesions produce little or none, unless epileptogenic activity develops with the generation of spikes, sharp waves, spike-wave complexes, etc.²

EFFECTS ON HEALTHY AGEING

The more important changes that accompany healthy ageing include alpha rhythm frequency slowing, changes in beta (fast) rhythm abundance, the appearance of diffuse delta activity, local anterior temporal delta wave foci and alterations in nocturnal sleep patterns^{3,4}.

Mean Alpha Rhythm Frequency

In young adults, this is around 10 Hz and does not change until 60 years. Subsequently there is a gradual decline, the rate of decrement varying from 0.05 to 0.75 Hz/decade across studies; 9.0–9.5 Hz at 70 and 8.5–9.0 Hz after 80 years. Indeed, in healthy subjects of any age it is rare to have a dominant (alpha) rhythm frequency of less than 8 Hz. In elderly patients, the degree of alpha frequency slowing is significantly related to the extent of cognitive impairment.

Low-voltage Beta (Fast) Activity

This is a low voltage 15–30 Hz rhythm, arising from the frontocentral areas of both hemispheres and sometimes diffuse in distribution, which increases in abundance in middle age and is more prominent in females. It persists into old age but diminishes markedly after 80 years.

Diffuse Slow Activity

This consists of background theta (4–7 Hz) and delta (1–3 Hz) waves of generalized distribution, is not seen in healthy people under 65 years and is relatively rare in early senescence (7% of people under 75 years), but occurs in up to 20% of persons over 75 years. Even in this relatively aged population the degree of diffuse slowing is mild. When moderate or severe, there is a significant association with a clinical diagnosis of dementia.

Focal Delta Waves

A substantial minority (30–40%) of healthy people over 60 years have focal delta (slow) waves strictly localized to the anterior temporal region and often lateralized to the left side. More extensive spread beyond the anterior temporal area indicates local pathology.

Paroxysmal Activity

Paroxysmal EEG events are transient, higher voltage waveforms that arise suddenly from the EEG background. The healthy brain generates some normal paroxysmal activity, namely lambda waves during wakefulness when the eyes are open and paroxysmal waveforms during sleep as part of the alerting process (lambdoid waves in light sleep and K-complexes in medium to deep sleep). Abnormal paroxysmal discharges include bifrontal delta wave episodes, spikes, and sharp waves and spike-wave complexes. Such phenomena are extremely rare in healthy elderly people.

Nocturnal Sleep Patterns

EEG sleep studies of healthy elderly individuals show decreases in sleep efficiency, greater numbers of awakenings from sleep with increases in total time awake (especially in the last 2 h of the night) as well as marked diminution in stages 3 and 4 sleep. These

changes, which usually begin somewhere around 50 years of age, are age-related and generally more pronounced in men⁵. A shortened rapid eye movement (REM) latency may be helpful in separating early dementia (normal REM latency) from depression (shortened REM latency). The former show decreased amounts of sleep spindles and K-complexes with reduced REM sleep percentage and a normal REM temporal distribution.

EEG IN DEMENTIA

With important exceptions, the changes in dementia are qualitatively similar to those of healthy ageing, although the degree of change is much more marked. Since the days of Berger, there have been many studies of the routine clinical EEG in dementia (reviewed by Busse⁶, Pedley and Miller⁷, Fenton⁴).

Changes Common to Most Dementing Disorders

Slowing of the dominant, parieto-occipital (alpha 8–13 Hz) rhythm over both hemispheres, a moderate to marked increase in generalized theta (4–7 Hz) and delta (1–3 Hz) activity (diffuse slowing) and a bilaterally symmetrical decline in low voltage beta (fast) activity are common background activity changes. One autopsy study reports a significant correlation between alpha frequency slowing and the number of senile plaques counted in Alzheimer's disease (AD) patients' brains⁸. Paroxysmal runs of bifrontal delta activity are not uncommon in dementia patients. In one investigation these have been related to degenerative brain stem changes at autopsy⁹. The occipital responses to photic stimulation at fast flicker rates (equal to or greater than 18 flashes/s) tend to disappear in a significant minority of dementia patients (1 in 5).

Differences Between the Various Dementias

AD vs. Pick's Disease

Studies that have compared the various types of dementing illnesses indicate that less than 5% of patients with histologically confirmed AD have a normal EEG even when first referred to the psychiatric services^{4,10–12}. In contrast, Pick's disease and multi-infarct dementia (MID) are not infrequently associated with normal EEGs. The number of Pick's disease patients in any study is small, but a consistent finding is that around 50% have normal records. Even when diffuse slowing is present in Pick's disease patients, the alpha rhythm is better preserved^{9,12,13}. The alpha rhythm is generated by the parieto-occipital areas of the cerebral cortex modulated by thalamocortical influences. The histological changes in Pick's disease are largely confined to the frontotemporal regions, which are relatively "silent" electrically, compared to the parieto-occipital areas, where the predominant AD changes occur. Indeed, it has been recently suggested that a normal EEG is one of the characteristic features of dementia of frontal lobe type: a dementing syndrome with onset in the presenium and selective frontal lobe dysfunction. It is not clear whether it represents a form of Pick's disease¹⁴.

Multi-infarct Dementia (MID)

The EEG in MID differs from AD in displaying significantly more asymmetry between the hemispheres, localized slow wave disturbances being particularly common, while the alpha tends to be better preserved. For example, Constantinidis *et al.*¹⁵ report three

times more alpha rhythm and five times more local slow wave foci in MID. Often, the laterality of the EEG focus in MID correlates with past or present clinical evidence of an ischaemic lesion lateralized to the same side. In contrast, AD patients have a significantly higher incidence of diffuse delta activity.

Huntington's Disease

The EEG in Huntington's disease differs from the dementias already discussed. In a variable number (30–80%) depending on the series reported, a low voltage tracing with an average amplitude of 10 μ V or less is a characteristic feature. This amplitude reduction correlates with caudate nucleus involvement but only becomes apparent by the time the disease is clinically well-established^{16,17}.

The Significance of Paroxysmal Abnormalities with Periodicity

Paroxysmal bifrontal runs of delta waves are common in dementia patients, especially those with AD. In one histological investigation, this bifrontal delta activity has been related to degenerative brain stem changes⁹. Regularly recurring (periodic) generalized biphasic or triphasic sharp wave or slow wave complexes of generalized origin with a characteristic recurrence rate of 0.5–1.0 s (intervals between successive bursts of complexes) are a characteristic feature of Creutzfeldt–Jakob disease (CJD). Early in the illness, diffuse background slowing occurs and in most cases the characteristic periodic complexes emerge. In a minority of patients (up to one-third in some series), especially the amyotrophic cases, this pattern may not be seen or may appear late. If practical, serial recordings are recommended¹⁸. On rare occasions, the periodic discharges may be temporarily focal, later generalizing and becoming bilaterally synchronous as the disease progresses. Their presence in a middle-aged or elderly patient with dementia is highly suggestive of CJD. Periodic triphasic waves of generalized distribution can be seen in other conditions, notably hepatic and other metabolic encephalopathies, subacute sclerosing leucoencephalitis and Unverricht's myoclonus epilepsy. Rarely they may appear in advanced AD patients and in Binswanger subcortical encephalopathy but do not show the characteristic periodicity or evolution of CJD.

EEG, COGNITIVE AND CT SCAN CHANGES

McAdam and Robinson¹⁹ reported a correlation of +0.79 between ratings of EEG change and clinical severity in dementia patients of mixed aetiology. The association between EEG slowing and severity of dementia has been replicated by many subsequent studies^{4,6,20}. However, Johannesson *et al.*¹² report that this electroclinical relationship held for 100% of their AD cases but almost half of their Pick's and MID patients had normal EEGs in the presence of significant cognitive decline. The correlation between the EEG slowing and extent of cortical atrophy as measured by computed tomography (CT) is weak, the link being obvious only in advanced cases. The EEG correlates better with clinical scales sensitive to early dementia, while the converse is true for the CT scan²⁰. Combining the two measures improves their diagnostic power. A discriminant function analysis study of 56 AD patients and 84 normal controls correctly classified 86% using the EEG data and 84% using the CT scan information. Combining the EEG and CT scan variables improved the correct classification rate to 90%². Interestingly enough, the degree of functional brain impairment as measured by

the EEG predicts survival time while the extent of cortical activity does not²².

THE APPLICATION OF QUANTITATIVE ELECTROENCEPHALOGRAPHY

Recent computerized EEG (CEEG) investigations have used frequency spectral analyses of the background activity and have generally replicated the earlier reports obtained by visual inspection of the EEG tracings. An overall slowing in mean EEG frequency is an invariable finding, with relative power increases in the slower frequencies (theta/delta) and decreases in the fast (beta) frequencies. The degree of frequency slowing correlates reasonably well with clinical ratings of dementia severity, e.g. Mini-Mental State and Clinical Dementia rating scales.

Is there a Relationship to Clinical Severity?

Alterations in the distribution of power across the various frequency bands seem to relate to the severity of the dementing process. In less severe cases, the main changes are increases in theta and reductions in beta power. When the disease is advanced, the respective amounts of delta and alpha power are also affected, the former being increased and the latter reduced. These associations have been observed during cross-sectional studies, there being a paucity of longitudinal investigations.

What About Early Cases?

Much of the work has been on populations of patients with established AD being cared for in hospital. A number of recent studies have examined mild probable AD patients (clinical dementia ratings of 1) compared to age-matched controls. These have replicated the earlier findings of frequency spectral slowing which have a specificity of virtually 100%²³. However, the sensitivity is only 20%. This means that only one in five of early cases of AD will have CEEG findings that deviate three standard deviations from the mean and therefore fall into the AD range.

The Potential Role of Brain Mapping

A new development in EEG technology has been brain electrical activity mapping (BEAM). This involves sampling the cortical electrical activity from multiple scalp electrodes and calculating the distribution of voltage across the whole of the scalp using a mathematical interpolation method. The patterns of scalp voltage distribution are displayed as colour-coded contour maps. This technique investigates the patterns of electrical activity generated at the same time by different areas of cerebral cortex. It can be used to detect regional differences and promises to be a useful tool in the investigation of the dementias. Several investigators have used multichannel or BEAM recordings to demonstrate localized left temporal lobe delta power abnormalities, which may in AD patients be an early manifestation of the disease²⁴⁻²⁷. Hence, BEAM may prove especially useful in the early detection of dementia.

Does the CEEG Change as the Dementia Progresses?

The few longitudinal CEEG studies that have been carried out over several years give conflicting results. Some report significant

decreases in mean frequency over time, with increases in delta/theta power and reduced alpha and beta activity²⁸⁻²⁹, but others have found either statistically insignificant trends in the direction of frequency slowing or that only about half of AD patients show progressive EEG changes over 12 months³⁰⁻³¹. It is noteworthy that the negative reports deal with relatively short time scales; 18 months or less. This may not be long enough to establish significant progression. There is also some evidence that frontal lobe quantitative changes may precede more generalized slowing³¹.

Differences between AD and MID

Few CEEG investigations have investigated the question of quantitative differences between AD and MID patients. A recently completed study in my laboratory reveals significantly greater amounts of delta power and less alpha power in the temporal and parieto-occipital areas of both hemispheres in AD patients, compared to those with a clinical diagnosis of multi-infarct dementia. As well as having less alpha power, the peak alpha frequency is slower in AD patients with mean values of 7 Hz and 8 Hz in AD and MID subjects, respectively, and significant asymmetry between the hemispheres in the MID patients³¹.

The degrees of synchrony between different areas of cortex within each cerebral hemisphere and between homologous areas of the right and left hemisphere can be assessed by coherence spectral analyses, which measures the similarity between pairs of EEG signals generated by different cortical areas. The coherence function is essentially a frequency correlation coefficient and measures the correlation at each frequency. It varies from 0 (signals quite different) to +1.0 (signals identical).

The main coherence differences between the AD and MID patient are seen in the temporal and parieto-occipital areas of both hemispheres. Compared to the MID patients, the within-hemisphere alpha and beta synchrony is lower in AD subjects. The pattern of between-hemisphere synchrony is different, being higher in AD patients for theta components between the temporal areas and lower for the alpha and beta frequencies between the parieto-occipital areas³¹.

Do Elderly Patients with "Non-organic" Psychiatric Illness Deviate from Normals?

In my laboratory we have also investigated age-matched controls and elderly patients with major depressive illness. Compared to the normals, the depressive patients had significantly more theta power and less alpha power, such changes being maximal in the temporal regions. Indeed, the mean spectral values of the three patient groups could be ranked roughly according to degree of deviation from the healthy controls. The AD patients were the most deviant, then those with MID, and finally the depressed patients. The deviation of the elderly depressives from the healthy controls raises the issue of the contribution of organic brain disease to the genesis of affective disorder in old age, as suggested by current CT and single photon emission computed tomography (SPECT) scan work^{32,33}.

EVENT-RELATED POTENTIALS IN DEMENTIA

What is an Event-related Potential?

Event-related or evoked potentials (ERPs or EPs) consist of transient voltage changes that occur in response to a sensory stimulus. These take the form of a series of negative and positive

waves, which last a number of milliseconds (ms) and measure a few microvolts (μV) in amplitude. They are "buried" amongst the "noise" of the ongoing EEG and are "extracted" by summing or averaging the response to a series of identical stimuli. This results in a waveform that lasts from a few ms up to several s.

Types of Event-related Potential

It takes about 20 ms for information to reach the cerebral cortex from a peripheral sense organ. Hence the early ERP components reflect neuronal activity in the sensory receptor itself and the afferent pathways of the brainstem. They depend on the functional integrity of the relevant sensory system, being relatively impervious to changes in psychological state. The brainstem responses consist of five positive (I–V) waves within 10 ms of the stimulus, and are especially stable and stimulus-bound, having prolonged latencies in brainstem disease. In contrast, the middle-latency (80–200 ms) and long-latency (> 200 ms) components are influenced by attention processes and how the subject perceives or processes the stimulus. The long-latency waves are termed cognitive or endogenous potentials, since their form is largely determined by the subjects' psychological state.

Healthy Ageing

The latency and waveform of sensory ERPs are critically affected by peripheral receptor changes. Hence, impaired hearing and visual acuity, common in the elderly, can cause problems in interpretation. If such sensory deficits are controlled for, the early and mid-latency potentials are little affected by healthy ageing. In contrast, the P_{300} wave, a positive going cognitive ERP that appears about 300 ms after a subject receives an important but unexpected stimulus, declines in latency and amplitude with advancing years.

Brainstem ERPs

Since there are brainstem changes, neuronal cell loss and neurofibrillary tangles in AD early ERP abnormalities can be predicted. The available brainstem auditory evoked response data are conflicting, two studies reporting delayed central conduction times (prolonged I–V intervals) with negative findings in a third^{34–36}.

Visual ERPs

A promising finding is the differential latency pattern of visual evoked potentials in dementia. Wright *et al.*³⁷ report completely normal pattern reversal responses with significant latency delay of the major positive component (P_2) of the flash response. The difference value between the prolonged flash P_2 latency and the normal latency value of the equivalent positive (P_{100}) wave of the pattern response is much longer in dementia patients compared to controls. It has the advantage of being little affected by drugs and only slightly lengthened by the ageing process, but does not discriminate between AD and MID³⁸. These findings have been replicated twice^{39,40}.

Cognitive ERPs

The P_{300} wave is elicited by a task requiring discrimination between two types of stimuli, one frequent and the other rare in

repetition. It develops as a response to the rare stimulus, which is used as a target and is a measure of cognitive processing time. Its latency may well reflect the time required for stimulus evaluation and categorization, while the amplitude is inversely related to probability of occurrence of the target stimuli. Healthy ageing increases the latency by around 1.36 ms/year and decreases the amplitude at a rate of about 0.18 $\mu\text{V}/\text{year}$ ⁴¹. The P_{300} latency is prolonged in many dementia patients by more than 1.5 standard deviations from the normal age-related mean and lengthens progressively over time^{42,43}. Positron emission tomography (PET) scanning of early patients has shown that P_{300} latency is inversely correlated with the relative metabolic rates of the parietal and, to a lesser extent, temporal and frontal association areas, but not with the subcortical areas⁴⁴. There is also a significant positive correlation between P_{300} amplitude and CSF 5-hydroxyindoleacetic acid (5-HIAA) concentration in AD patients⁴⁵.

Distinctions between Cortical and Subcortical Dementia

Some work suggests that auditory ERP patterns may be useful in distinguishing cortical from subcortical dementia. The P_{300} wave is abnormal in both. In contrast, the earlier waves (middle-latency components; N_1 , P_2 and N_2) are intact in the former but abnormal in the latter⁴⁶. Indeed, a recent preliminary study reports correlations with specific aspects of cognitive functioning; N_1 and P_2 with motor speed and N_2 with short-term memory⁴⁷.

OVERVIEW

The potential application of the ERP recordings to the diagnosis of dementia has yet to be clearly defined. None of the available techniques are capable of distinguishing between AD and MID. The visual flash-pattern reversal latency pattern is a robust finding in established cases of dementia of cortical origin. Further, there is some evidence that it reflects a specific cholinergic deficit. However, the abnormality rate in patients during the early stages of dementia remains to be determined. On the other hand, the P_{300} latency delay has been shown to be present in a majority of recent-onset AD patients⁴⁴. A difficulty is that the correct classification rate for AD patients using the P_{300} data alone has varied widely across studies, from 89% to 20%, most having rates of more than 70%. This variability reflects not only differences in the clinical material and selection of controls but also the type and level of task difficulty of the experimental paradigm used to record the P_{300} wave. The paradigm has differed in the various studies. Research is required to determine the optimum paradigm for eliciting abnormality in early dementia cases, so that standardization across studies becomes possible. Further work on central brainstem conduction times and the pattern of middle latency components in subcortical dementia is also indicated.

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Computed Tomography (CT)

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Computed tomography (CT) was introduced in the early 1970s and has become one of the standard investigations in the clinical neurosciences. In cranial CT, X-radiation is passed through the head in the form of a tightly collimated beam and is measured by a series of detectors. Radiation is absorbed by the intervening structures, which results in “attenuation” of the beam. Attenuation is maximal in high-density regions such as bone and minimal in low-density regions such as cerebrospinal fluid (CSF). The information from the detectors is processed by computer and the product is a numerical output. The brain is divided into three-dimensional volume elements (or voxels) and each is given an attenuation number, which represents the average attenuation in that area. The numerical output is transferred to a grey scale and each voxel is represented by a two-dimensional pixel. The familiar CT scan images are the result of the pictorial representation of the pixels on the grey scale. Areas of high attenuation are represented by white and areas of low attenuation by black. Areas with intermediate attenuation (such as brain substance) appear grey. The amount of radiation exposure in an average CT scan is slightly less than that in a set of conventional skull X-rays.

USE OF CT IN CLINICAL PRACTICE

CT has several advantages over magnetic resonance imaging (MRI) in the investigation of dementia. Acute haematomas can be easily distinguished from areas of infarction, the increased density of the former contrasting significantly with the hypodensity of the latter. In practice, CT is more widely available than MRI and the examination is less arduous and can be completed much faster. The presence of a cardiac pacemaker or the presence of surgical clips from previous brain surgery are not contra-indications to CT scanning in the same way that they are for MRI.

Guidelines have been published that suggest the circumstances under which a CT scan should be performed for the work-up of dementia^{1,2}. Essentially, where the duration of the illness is short (<6 months and certainly <3 months), and where the features of the illness indicate that there may be cerebral pathology, then the chances of a CT scan detecting a clinically significant lesion is increased. Such features include: focal neurological signs; epileptic fits; variations in the course of the illness; and indicators of the presence of normal pressure hydrocephalus (gait disturbance, and incontinence in the presence of dementia). One study specifically examined a population sample aged 65+ and found that potentially treatable lesions (subdural haematoma, hydrocephalus, non-metastatic intracranial tumour) were present in 145 out of a possible 137000 patient years at risk³³. Specific features predicting the detection of such a lesion were: cognitive

impairment for 1 month or less; head trauma in the week before mental state change; rapid onset of change over 48 h; history of CVA; seizures or incontinence; focal neurological signs; papilloedema; visual field defects; gait abnormalities; postural instability; or headaches. Paris *et al.*³ estimate that the yield of potentially treatable conditions is about 3%. Factors on CT scan that predict who will respond positively to a CSF shunt have been documented by Vanneste *et al.*⁴—cerebral atrophy and the presence of white matter disease were poor predictors of response to the insertion of a shunt.

DIFFERENTIAL DIAGNOSIS

CT scanning is helpful in confirming certain diagnoses, e.g. Alzheimer's disease (AD), but is not the definitive investigation, whereas in other disorders, e.g. 'subdural haematoma' the CT is the definitive investigation.

The CT scan has been used to differentiate vascular dementia from primary degenerative dementia. Generally speaking, good concordance between the presence of vascular lesions on CT and the presence of vascular dementia (defined purely clinically or using the Hachinski score) has been achieved⁵⁻⁷.

White matter lesions on CT scan have been widely reported^{8,9} and are associated with impaired cognitive function (in both AD patients and non-demented subjects) and neurological signs (gait disturbance and extensor plantar response). Scheltens *et al.*¹⁰ described a number of rating scales used to detect white matter changes on CT and MRI brain imaging, concluding that the ideal rating scale is not yet in existence but that different rating scales serve individual purposes.

Excessive ingestion of alcohol can result in cerebral atrophy and ventricular dilatation on CT scan, particularly affecting the frontal lobe and cerebellar vermis. It is apparent that the changes occur relatively early (but do not antedate alcohol excess), are apparent before any clinical evidence of declining cognitive function and may be partially reversible with abstinence¹¹. There is also evidence that third ventricular size is correlated with memory impairment in alcoholics without Korsakoff's deterioration. Patients with Korsakoff's psychosis have more cortical atrophy and lateral ventricular enlargement, but the size of the third ventricle is particularly increased.

Depression has been shown to be accompanied by both cerebral atrophy and ventricular enlargement^{12,13}. CT scan appearances in depressed patients appear to be midway between those of normal controls and demented subjects, tending to be nearer the latter. More recently, it has been shown that patients with reversible dementia secondary to depression (pseudodementia, or dementia

Table 63.1 Clinical and structural correlates on CT in AD

Reference	Mean age (or range)	n	CT measure	Clinical features	Association†
36	70.0	43	Cortex	GRD	C=0.56
			Lateral ventricles		C=0.62
37	78.6	40	Cortex	Age, paranoid delusions	NS, C = -0.65**
			Lateral ventricles	Age, digit symbol test	NS, C = -0.31*
			Evan's ratio	Age, digit symbol test	NS, C = -0.31*
38	NS	22	Lateral ventricles	GRD, duration	C=0.73**_NS
			Sylvian fissures	GRD, duration	C=0.59*, NS
			Surface sulci	GRD, duration	NS, C=0.61**
			Cortex	Age	NS
(a) Presenile	59.6	10	Cortex	GRD, age	NS, C=0.27*
(b) Senile	77.6	7	Lateral ventricles	GRD, age	C=0.29*, C=0.50***, C=0.46***
40	53–87	59	III ventricle	GRD	NS
			IV ventricle	GRD	NS
41	77.0	57	Cortex	Age, duration, cognition, AD	NS, NS, NS, NS
			Lateral ventricles	Age, duration, cognition, AD	NS, C=0.29*, C=-0.35**, C=-0.43***
			III ventricle	Age, duration, cognition, AD	NS, NS, C=-0.40***, C=-0.30**
42	72.7	35	Lateral ventricles	Age, GDS, MSQ	C=0.55*, C=0.37*, C=0.42**
32	58.1	8	Lateral ventricles	Memory	C=-0.68**
			Lateral ventricles	Verbal fluency	C=-0.59**
			Lateral ventricles	Proverb interpretation	C=0.74***
			Lateral ventricles	Clock drawing	C=0.60**
43	78.1	47	Lateral ventricles	Digit copying test, MTS	C=0.24**, NS
			Regional density: pontine		C=-0.31***, C=0.40**
44	60.7	60	Cortex	Aphasia, GRD	NS, NS
			Lateral ventricles	Aphasia, GRD	*, *
45	67.9	42	Cortex	Memory, deterioration in IQ	C+0.30*, NS
			Lateral ventricles	Memory, deterioration in IQ	NS, NS
46	45–84	39	Total intracranial CSF	IMC, MMSE, BDS	*, *
47	72.2	16	Lateral ventricles	MMSE	C=-0.46*
31	63.4	30	Cortex	Age, duration, MMSE, GDS	NS, *, ***, ***, **
			Lateral ventricles	Age, duration, MMSE, GDS	NS, NS, NS, NS
			III ventricle	Age, duration, MMSE, GDS	**, NS, ***, ***, **
17	79.7	138	Cortex	Age, duration, MMSE, CAMCOG	C=-0.27***, C=0.24**, C=-0.42***, C=-0.41***
			Lateral ventricles	Age, duration, MMSE, CAMCOG	NS, NS, C=-0.25**, C=-0.31***
			III ventricle	Age, duration, MMSE, CAMCOG	NS, C=0.22**, C=-0.31***, C=-0.34***
			R Sylvian fissure	Age, duration, MMSE, CAMCOG	C=0.21**_NS, C=-0.25**, C=-0.27**
			L Sylvian fissure	Age, duration, MMSE, CAMCOG	NS, NS, C=-0.28***, C=-0.30***
48	79.7	138	Lateral ventricles	Delusions	** (Smaller ventricles)
			Basal ganglia calcification	Delusions	**
			Temporal atrophy	Delusions	*
			Frontal/occipital atrophy	Aggression	***
50	75.0	60	Brain quadrants	Hyperorality	***
				Misidentification symptoms (MS)	MS associated with larger right anterior horn of lateral ventricles and left anterior brain areas
51	75.0	60	III ventricle	Age, duration, MMSE, CAMCOG	NS, C=0.34**_C=-0.23**_C=-0.27**
			L anterior horn	Age, duration, MMSE, CAMCOG	NS, C=0.39***, C=-0.33***, C=-0.36***
			R anterior horn	Age, duration, MMSE, CAMCOG	NS, NS, C=-0.25**, C=-0.23**
			Subarachnoid areas:		
			L frontal	Age, duration, MMSE, CAMCOG	NS, C=0.32**_NS, NS
			R frontal	Age, duration, MMSE, CAMCOG	NS, C=0.39***, NS, NS
			L posterior	Age, duration, MMSE, CAMCOG	NS, C=0.36**_NS, NS
			R posterior	Age, duration, MMSE, CAMCOG	NS, NS, NS, NS
			Total intracranial density	Age, duration, MMSE, CAMCOG	NS, NS, NS, NS
52	75.0	60	Grey matter	Age, duration, MMSE, CAMCOG	NS, NS, NS, NS
			White matter	Age, duration, MMSE, CAMCOG	NS, C=0.32**/_NS, NS

†Associations reflect the information in the original paper: NS, not significant; C, correlation coefficient (Spearman or Pearson), given if significant followed by significance level: * $p < 0.05$, ** $p < 0.01$; *** $p < 0.001$.

/In L parietal only.

GRD, Global rating of dementia (varies from study to study); GDS, Global Deterioration Scale; MSQ, Mental Status Questionnaire; MTS, Mental Test Score; MMSE, Mini-Mental State Examination; CAMCOG, part of Camdex³; IMC, Information memory and Concentration Test; BDS, Blessed Dementia Scale. Table reproduced from O'Brien, J., Ames, D., and Burns, A. (eds), *Dementia*, 2nd edn, 2000, by permission of Edward Arnold.

syndrome of depression) have CT scan changes such as increased lateral ventricular size and decreased tissue density numbers¹⁴. In paraphrenia, there is dilatation of the lateral ventricles, preservation of cortical structures and loss of the normal ventricular size/age correlation seen in normal ageing¹⁵.

CT IN AD

Two areas of clinical interest are important in relation to CT scanning in AD. First, what is the diagnostic ability of the CT scan? Second, what CT changes take place in AD and how are they related to the clinical features of the disorder?

DIAGNOSTIC ABILITY OF CT

The second area of clinical interest relates to the ability of CT scan changes to differentiate patients with dementia from non-demented control subjects. It is well recognized that demented subjects can have normal CT scans, whereas normal subjects can have marked atrophic changes. Discriminant analyses are able to differentiate the two groups using CT scan appearances in about 80% of cases¹⁶, a rate that has remained virtually constant over time in spite of advances in CT technology and methods of scan analysis⁴⁹. In a meta-analysis, De Carli *et al.*¹⁸ estimated sensitivity and specificity for a variety of CT measures. Specificity was high for most measures (about 90%, i.e. few normal subjects were classified as having abnormal CT appearances). Sensitivity (i.e. the number of AD patients regarded as having abnormal CT scans) was lower.

Serial CT scans in individual patients have been performed and have the potential for greater diagnostic accuracy. Increases in ventricular size have been shown to outstrip those which take place in normal ageing. Luxenberg *et al.*¹⁹ found that the rate of lateral ventricular enlargement in male AD patients over 12 months completely differentiated these patients from controls (i.e. 100% sensitivity and 100% specificity). Increase in ventricular size correlates with deterioration in cognitive function¹⁹ and two subgroups of patients with AD have been described on this basis of one with significantly increasing ventricular size and deteriorating cognitive function and one without these changes²⁰.

CLINICO-RADIOLOGICAL CORRELATIONS

Early studies demonstrated correlations between the degree of intellectual impairment and both cortical and subcortical atrophy on CT²¹⁻²³ but often included normal controls in the correlations. Normal control subjects tended to be patients referred for investigations and found to have normal scans, rather than people screened first and then scanned.

With regard to clinico-radiological changes, both cortical atrophy and lateral ventricular enlargement occur with normal ageing and tend to accelerate after the age of 60. There is a significant correlation between cerebral atrophy and age in normal subjects but this has not been as consistently found in dementia. Correlations have been described between cognitive function and both cortical atrophy and ventricular size. Generally, correlations are higher in the latter relationship (ventricular size can be measured as a continuous variable, which may partly explain the greater association), although some studies have reported no association between degree of dementia (measured by both specific cognitive tests and global ratings) and either CT measure. The third ventricle has been examined in a number of studies, and was found to be larger in demented patients than in

age-matched controls, correlating with degree of cognitive impairment^{17,24}.

In addition to measures of global cerebral atrophy and ventricular enlargement, CT scans can provide other information of clinical interest. Regional cerebral atrophy has been shown to be related to certain behavioural disturbances⁴⁹. Basal ganglia calcification is found in a significantly greater proportion of patients with delusions and demented patients with affective symptomatology have less severe CT scan changes, including relative preservation of the interhemispheric fissure⁴⁹.

Table 63.1 summarizes the relevant studies; 88% of ventricular measures show significant correlations with cognitive tests, whereas only 41% of cortical assessments do so ($\chi^2 = 11.3$, $p < 0.001$, d.f. = 1).

The diagnostic potential of the specific temporal lobe views of the brain has attracted some interest in AD²⁵. Pathologically, the temporal lobe discriminates well between AD and normal controls²⁶ and coronal plane CT images can be reformatted to display the temporal lobes in fine detail.

CT has been combined with SPET (single-photon emission tomography) in order to improve diagnostic accuracy in AD. Jobst and colleagues, in the OPTIMA project in Oxford^{27,28}, reported a series of studies demonstrating that views of the temporal lobe could be achieved during CT scan, with the plane orientated along the long axis of the medial temporal lobe (20–25° anterior to standard CT angle). In this way, 92% of patients with AD were correctly diagnosed compared to a 5% false-positive rate. Simple measurement of the narrowest thickness of the medial temporal lobe (right or left) was about 50% thinner in patients with AD compared to controls. Combining these measurements with SPET in patients with histologically proven AD compared to controls, the medial temporal lobe atrophy provides 94% sensitivity and 93% specificity, parietotemporal hypoperfusion on SPET gave 96% sensitivity and 89% specificity, and the combination of both changes gave a sensitivity of 97%. The results of Lavenu *et al.*²⁹ who carried out a similar study using CT and SPET, resulting in a diagnostic accuracy rate which was much less—68%. Stage of disease is an important influencing factor in this rate and replication of the results of these studies is needed before they can be incorporated into clinical practice. O'Brien *et al.*⁷ found reduced temporal lobe width in AD, vascular dementia and Lewy body dementia compared to controls, suggesting a lack of specificity of the finding with a single cross-sectional measurement.

CONCLUSION

In summary, the role of the CT scan in old age psychiatry is as a relatively non-invasive and widely available neuroradiological technique to exclude intracranial mass lesions. Regional changes may be helpful in the differential diagnosis of the dementia syndrome. The concordance between the CT changes and a clinical diagnosis of dementia is not absolute and significant overlap exists between the changes seen in dementia and those seen in normal ageing. Some methods of CT scan analysis are better than others in this differentiation and serial CT scans on individual patients may be an even better indicator.

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Magnetic Resonance Imaging

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The property that is the basis for magnetic resonance imaging is the interaction between hydrogen and a magnetic field. A hydrogen atom is a nuclear magnet which, when placed on the magnetic field, aligns itself in the direction of the field. In the process of aligning it develops a spin at a frequency which is called the Larmor frequency. If the magnetic field is varied, then the frequency of this oscillation also varies. Besides hydrogen, when a nucleus contains an odd number of either protons or neutrons, it is magnetic. As indicated earlier, the frequency of oscillation of hydrogen is proportional to the strength of the magnetic field. Resonance occurs when the applied frequency (magnetic frequency) is the same as that of the object. Magnetic nuclei in a magnetic field can be stimulated by other magnetic fields. These fields can usually be created by radio-frequency waves. When the radio-frequency waves are tuned to the Larmor frequency of the atomic nuclei, namely hydrogen, then the direction of oscillation of the atom moves towards the direction of the newly applied magnetic field. When the field is turned off, the nuclei relax, emit absorbed energy and return to the state prior to the radio frequency stimulation. This process can be repeated over and over again, provided that enough time is allowed for the relaxation of the atom. When consideration is given to an entire object composed of billions of atoms, the relaxation of these atoms will depend upon local effects, i.e. what molecule the atom is attached to, etc., the most common attachment being water for hydrogen. The signal detected when the hydrogen atom relaxes after the radio frequency and pulse has stopped can be detected by using a radio frequency antenna tuned to the Larmor frequency. The decay of this signal over time is called free induction decay. The time to recover two-thirds of the magnetization after stopping the radio-frequency stimulus is called the T1 relaxation time, or spin lattice relaxation time. This relaxation time is influenced by the environment of the hydrogen atom and can vary between gray matter, white matter, and fluid. T1 primarily refers to the longitudinal vector of relaxation, which is along the direction of the magnetic field. Another vector at right angles to the field, called the transverse vector, can also be measured, and its relaxation is called T2 relaxation. By convention, this refers to the time when it disappears, rather than the two-thirds used for measuring T1. T2 is always shorter than T1. Based on the properties of the longitudinal and transverse vector, one acquires images that are predominately T1-weighted, or those which are predominately T2-weighted, a third type that appear intermittent in appearance. In a T1-weighted image, fluid appears dark. In a T2-weighted image, fluid appears bright and white. Scanning sequences can be optimized by emphasizing one contrast vs. another. This allows a better distinction of both pathology and normal tissue.

MAGNETIC RESONANCE MORPHOMETRY

Magnetic resonance morphometry basically utilizes the acquired images to identify objects and then to quantitate their volumes. Volumes can be estimated in either two dimensions or three. The most common method of estimation is to use two-dimensional slices through a given object. A number of studies have indicated that systematic sampling is better than the randomized sampling of the particular object of interest. Segmentation of the object can be accomplished manually or by semi-automated means, and more recently in an automated fashion. Factors that effect the sensitivity and accuracy of the estimation of the volume include the number of slices that go through the object, the orientation of the slices, contrast, and any inhomogeneities in the magnetic field. Magnetic resonance morphometry methods have greatly improved over the last decade and are now widely utilized for studying a variety of neuropsychiatric disorders. The techniques, which were initially primarily manual outlining of an object, have vastly improved and now include semi-automated and automated techniques to estimate the volume of the object.

GERIATRIC PSYCHIATRY

The application of MRI in geriatric psychiatry has been extensive. There are two broad areas to which MR techniques have been applied. One is to evaluate the presence of gross pathology and the other is to evaluate morphometric changes in a variety of disorders in the elderly. MRI is often utilized to look for tumors, infarcts, etc. when one suspects disease in a patient. MRI can detect space-occupying lesions and, given that images can be acquired in three dimensions, it can often provide better resolution than computed tomography (CT) (it must be kept in mind that newer forms of CT provide sufficiently high resolution). MRI is not useful in identifying calcifying objects, and in this particular case CT is far better. The use of contrast agents, such as gadolinium, can distinguish any breaking of the blood-brain barrier and often these agents are utilized to produce additional contrast of pathological tissue. Full-blown infarcts can be easily identified on MRI. In addition, MRI can be utilized to measure blood flow through large vessels, a technique known as called magnetic resonance angiography.

BRAIN ATROPHY

The second aspect as it relates to geropsychiatry is a measurement of brain atrophy and dementia. Patients with frontal temporal

dementia have significant atrophy of the frontal lobes. This can be easily visualized using MRI and quantitated if need be. Patients who have significant Alzheimer's disease (AD) will demonstrate observable hippocampal atrophy. Over time this can be measured and utilized for assessing the progression of the disease. MRI can be acquired with a different contrast, as noted earlier, and in three dimensions. Utilizing this one can develop specific methods to identify objects of interest and quantitate these objects. Fluid measurements of cerebrospinal fluid in different locations can also be easily measured. The technique also seems to be of particular use in mild cognitive impairment, in which commencing atrophy, especially of the hippocampus and temporal cortex, are often seen.

SILENT INFARCTS

Another area of growing interest in geropsychiatry is the identification of silent infarct or leukoencephalopathy. These are a function of aging. The risk factors are very similar to those of stroke, including high blood pressure, diabetes, etc. These changes are usually present in the deep white matter, the periventricular region, and subcortical nuclei such as the caudate putamen and thalamus. These silent strokes occur at the ends of the perforating

blood vessels of the brain. These blood vessels do not have collaterals and are therefore more susceptible to the development of occlusions and the ischaemia that results from them. These silent strokes have been characterized pathologically as areas of myelin pallor, true infarcts (dead tissue and lacuna). A lacuna is distinguished by the presence of fluid inside the area and this can be determined by a bright signal on T2 and a dark signal on the T1 images. These changes have been related to depression, bipolar disorder and dementia. They are present in both AD and vascular dementia. The contributing role of these factors in AD is controversial. Epidemiological studies indicate that the presence and extent of these changes is a good indicator of a development of dementia and it has the same odds ratio as APO-E studies.

In summary, MRI is widely utilized to study various types of neuropsychiatric disorders. It has particular application and utility for identifying pathology and atrophy in the context of dementia, and lesions such as silent strokes in late-life depression and late-life bipolar disorder.

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Functional Magnetic Resonance Imaging (fMRI)

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Functional MRI (fMRI) is a new technique that has allowed the non-invasive assessment of physiological function of the brain in humans. The phenomena of nuclear magnetic resonance has been utilized to accomplish this. The first attempt at producing a functional image of the human brain utilized an invasive technique of evaluating blood flow changes in the brain, assessed by using a contrast agent. The non-invasive method was introduced by Ogawa, and is based on a blood oxygenation level-dependent contrast (BOLD). This contrast originates from the homogeneity induced by deoxyhaemoglobin in red blood cells. Signal intensities in MRIs are therefore acquired in a manner sensitive to this BOLD contrast, which is effected with regional deoxyhaemoglobin content. The rationale underlying the changes in deoxyhaemoglobin is that regional blood flow increases, while the oxygen consumption rate is not altered significantly, resulting in a lower deoxyhaemoglobin content per unit volume of brain tissue. The signal intensity in the image acquired sensitive to BOLD therefore increases in active regions relative to those regions which are not active.

Another technique that has been developed, called EPISTAR and FAIR, utilizes tagging of blood spins to measure cerebral blood flow changes. In a recent study, we have demonstrated that, using EPISTAR, there is a reduction in blood flow in the left frontal lobe in depressed patients. BOLD as well as FAIR can be developed and utilized in regular MRI 1.5 tesla scanners. However, many particularly exciting findings are emerging from high-field MRI, 4 tesla and greater. BOLD response is greater in a higher magnetic field, which produces an improved contrast.

APPLICATION OF fMRI

fMRI has been mostly utilized with the BOLD type of measure to evaluate brain regions involved in a task performance. Unlike positron emission tomography (PET), fMRI allows a single trial design and a single subject design and measures images that can be acquired in tenths of milliseconds, which is still much slower than the temporal response of neurons but similar to the response of the vascular system. In the design paradigms in general, the contrast is measured before and during a task and this can be done repeatedly to obtain an average within a subject on/off task. It can also be used to track the evolution of a particular task over time, that is the temporal evolution of the signal relative to the execution of this task can be obtained and averaged following repeated executions of the same task. It also allows evaluation of learning, errors, habituation, etc. The technique has had limited applicability so far to disease conditions, except as it relates to the evaluation of working memory tasks and other components of memory in patients with mild cognitive impairment, as a prelude to assessing whether they are likely to develop dementia over time. Clearly, this technique offers a significant possibility for understanding many of the neuropsychological functions that are altered in aging and acquiring the ability to assess their changes relative to particular diseases of aging.

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Positron Emission Tomography (PET)

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Positron emission tomography (PET) provides quantitative images of the function of the brain in life. It generates an image of the distribution of a radioactively-labelled tracer substance that is distributed in the brain according to its pattern of physiological activity. By appropriate choice of tracer substances, it is possible to measure physiological variables such as blood flow, metabolism, neurotransmitter receptors, presynaptic neurotransmitter pools and aspects of amino acid metabolism.

The essence of any tomographic technique is a mathematical reconstruction of a two-dimensional image of the distribution of some physical variable (e.g. concentration of radioactivity), from a set of measurements of the value of that physical variable, averaged along a large number of intersecting straight-line paths through the brain. The feature that distinguishes PET from single photon emission computed tomography (SPECT) is the fact that a PET camera detects the paired photons generated by positron annihilation. When a positron is emitted from the tracer substance, it is annihilated by collision with an electron in the surrounding matter. This annihilation generates two γ -ray photons, which each have an energy of 511 keV and must travel in opposite directions. When two photons are detected simultaneously in different crystals arranged in a ring around the head, it can be concluded that a positron annihilation event has occurred at some point along the straight line connecting the two detectors. This use of coincidence detection of pairs of photons to determine the direction of travel of the photons is intrinsically more efficient than the use of collimators to determine the direction of travel, as is necessary in SPECT. Furthermore, correction for absorption in the brain tissue is straightforward in PET studies, because the amount of absorption along the straight-line path between a pair of detectors can be measured directly in a preliminary transmission scan, using a ring source. It is thus possible to quantify the local concentration of the positron-emitting isotope in the brain in absolute terms.

The positron-emitting isotopes used in PET are ¹⁵O [half-life for radioactive decay (T/2) 2 min]; ¹³N (T/2, 10 min); ¹¹C (T/2, 20 min); and ¹⁸F (T/2, 110 min). The relatively short half-life of positron-emitting isotopes presents a logistic problem, since the PET camera must be located near to the cyclotron required to produce the isotopes. However, an advantage of short-lived isotopes is that the rapid decay of radiation minimizes unnecessary radiation exposure after the procedure is completed. In addition, in the case of ¹⁵O, it is possible to carry out multiple separate PET scans within a single session of investigation, allowing the measurement of changes in brain

function in response to various mental or pharmacological stimuli.

PET TRACER SUBSTANCES

In principle, virtually any substance involved in physiological processes that can be labelled with a positron-emitting isotope might be used in PET, but quantitative measurement requires an adequate mathematical model of the processes governing distribution of the substance in the brain.

An image of regional cerebral blood flow (rCBF) can be obtained from the distribution of intravenous [¹⁵O]H₂O which is delivered to the brain at a rate depending on cerebral perfusion. Inhaled [¹⁵O]oxygen provides an image of oxygen distribution in brain tissue which, when combined with an rCBF image, can be used to generate an image of regional oxygen metabolism (rCMRO₂). Intravenous [¹⁸F]deoxyglucose (FDG) can be used to provide an image of regional glucose metabolism (rCMRGlu). This technique relies upon the fact that deoxyglucose is transported into cells by the same mechanism as is responsible for glucose uptake, but its metabolism is arrested after the first reaction in the glycolytic pathway. Hence, deoxyglucose accumulates within cells at a rate which reflects the rate of entry of glucose into the glycolytic pathway.

The development of labelled ligands that are suitable for measuring the characteristics of neurotransmitter binding sites is a difficult task, because of the difficulty in achieving a high level of specific binding at tolerable doses of the ligand. An example of a well-behaved ligand is [¹¹C]raclopride, which binds to D2 dopamine receptors¹. It associates and dissociates rapidly from receptors, so that equilibrium is established relatively quickly, making it possible to determine an equilibrium binding curve. From such a binding curve, it is possible to determine the strength of binding (KD) and the density of receptors (B_{max}). Other ligands that have been used to study neurotransmitter function include [¹¹C]SCH 23390 for D1 dopamine receptors², [¹¹C]WAY-100635³ to measure 5-HT₁ serotonin receptors, and [¹⁸F]altanserin⁴ and [¹⁸F]setoperone⁵ for measurement of 5-HT₂ receptors.

SOURCES OF VARIATION IN PET IMAGES

If an imaging technique is to be useful for delineating pathological processes, it is necessary that the variation in image due to the pathological process of interest should not be swamped by

variation arising from other factors. The difficulties in dealing with the multiple sources of confounding variation in PET images has hitherto limited the contribution of PET to the delineation of psychiatric disorders. From the clinician's viewpoint, the biological sources of variation are the most relevant, some of which are considered below.

In the study of elderly patients, one of the potentially confounding differences between subjects is brain atrophy, which is liable to produce variation in partial volume effects. Partial volume effects are due to the limited spatial resolving power of PET cameras, resulting in the contamination of the signal from high-intensity regions by adjacent tissue with low intensity. For example, if grey matter volume is reduced, the measured functional activity of the grey matter will be lowered by an increased contribution from adjacent white matter or CSF. A number of studies have examined the effect of cerebral atrophy on metabolic measures obtained with PET in both healthy older adults and patients with Alzheimer's disease^{6,7}. A recent study by Ibanez *et al.*⁸ used measures of grey matter volume obtained from MRI scans to correct PET FDG images in a group of mildly to moderately demented AD patients. They found that metabolic values in parietotemporal and frontal cortex were reduced compared to a control group, even after correction for atrophy. These data suggest that, despite the undoubted contribution of partial volume effects, the metabolic deficits seen in these patients are not due simply to atrophy, but reflect true decreases in metabolic activity.

When evaluating images of brain function, it is necessary not only to be aware of confounding variance introduced by extraneous factors, but also to know that the functional impairments in a single disease are quite heterogeneous. For example, the clinical manifestations of AD are diverse, including not only impairments of memory but also many different aspects of cognitive function and behaviour. In accord with this behavioural heterogeneity, PET studies reveal that cases differ in the degree of laterality of the abnormalities, and also in the degree of frontal involvement⁹. On the other hand, different diseases can produce phenomenologically similar mental states, raising the possibility that different diseases might produce similar patterns of perturbation of regional blood flow and metabolism. This possibility is illustrated by the fact that hypometabolism of frontal cortex has been reported in AD⁹, depression¹⁰, Parkinson's disease¹¹ and in the psychomotor poverty syndrome in schizophrenia¹². The greatest value of PET thus may lie in the sensitive measurement of particular patterns of brain malfunction, rather than the distinction between diseases differing in aetiology.

ACTIVATION STUDIES

A potentially useful approach to the measurement of brain function is within-subject comparison of brain activity during behavioural or pharmacological activation with that in an appropriate reference state. Behavioural activation has been used to identify the brain areas involved in specific mental activities in healthy young adults^{13,14} and in older individuals¹⁵. Interestingly, older adults have been found to have greater activation in frontal areas during some types of cognitive activity, compared to younger adults^{16,17}. These results have led to the suggestion that frontal cortex may play a compensatory role in the face of reduced function in other task-relevant brain areas, an idea that also is relevant to diseases of ageing.

Measurement of rCBF during pharmacological activation allows measurement of the response of the brain to drugs that alter neurotransmission. Although not yet used in psychiatric populations, the feasibility of this type of strategy has been demonstrated in normal subjects. For example, Friston *et al.*¹⁸

demonstrated that buspirone (a partial agonist at serotonergic 5-HT_{1A} receptors) caused an attenuation of the normal increase in rCBF in the parahippocampal gyrus produced by a verbal memory task, as well as a transient impairment of memory test performance. Conversely, Furey *et al.*¹⁹ have shown improved performance and reduced activity in the frontal lobe during a working memory task after administration of physostigmine, a cholinesterase antagonist. These studies suggest that modulation of neurotransmitter systems has specific effects on brain function during cognitive activity, and have interesting implications for the use of pharmacologic modulation in understanding psychiatric disorders.

BLOOD FLOW AND METABOLISM IN AD

PET studies have produced a moderately consistent picture of the abnormalities of blood flow and metabolism in AD. The most common pattern is bilateral reduction in parietal and temporoparietal flow and metabolism, especially in early cases^{9,20,21}, but the patterns of flow and metabolism show considerable variation between patients²². These variations correlate with variation in behavioural and neuropsychological impairments⁹. Longitudinal studies have demonstrated that focal metabolic abnormalities are detectable before the corresponding neuropsychological impairment becomes apparent, and that the metabolic abnormalities progress as the disease progresses²³. Recent studies have shown that metabolic abnormalities similar to those seen in the cortex of AD patients can be found in asymptomatic persons at risk for familial AD²⁴ and that these deficits are associated with the presence of the APOE-4 allele²⁵.

A few activation experiments with AD patients also have been conducted. Mildly demented AD patients have shown greater frontal activations, compared to healthy elderly, during memory tasks^{26,27} and during perceptual tasks²⁸. This is similar to the increased frontal activity seen in the healthy elderly, compared to young adults, and indicates that recruitment of cognitive resources mediated by frontal regions may be a common response to a decline in brain function, whether caused by normal ageing or by disease.

BASAL GANGLION FUNCTION IN HUNTINGTON'S DISEASE

The major neuropathological process in Huntington's disease (HD) is degeneration of the basal ganglia, especially the corpus striatum. Hence, it would be expected that there would be a loss of postsynaptic D1 and D2 dopamine receptors in the striatum, which has been amply demonstrated with imaging of these receptors using [¹¹C]SCH 23390 and [¹¹C]raclopride, respectively²⁹. It has been shown that a striatal decrease in dopamine is detectable even in asymptomatic persons who carry the HD gene mutation³⁰, and further that the degree of reduction is correlated with reduced performance on cognitive tests of frontal lobe function³¹. Glucose metabolism is also reduced in the striatum of HD patients and patients at risk for HD³². Longitudinal studies of disease progression in HD have shown declines in both striatum and frontal lobe metabolism³³, indicating that PET could be useful in following the effects of treatment over time.

Only a few activation studies have been carried out with HD patients. Patients have been studied with [¹⁵O]water during hand or finger movements and show reduced flow in the striatum, motor areas and prefrontal cortex^{34,35}. Of particular interest is that activity in parietal cortex in one study³⁴ was increased relative

to control subjects, suggesting a compensatory role of this region, similar to that postulated above for AD patients.

GERIATRIC DEPRESSION

Depression is a serious problem in older adults, and can occur as a symptom of a dementing illness or in the absence of dementia. The most common neuroimaging finding in depression is reduced glucose metabolism or flow in prefrontal cortex, although cingulate and paralimbic regions also are involved in mood disorders^{36,37}. These regional patterns of reduction are seen in elderly as well as younger patients, although elderly depressed individuals are more likely to have global metabolic reductions in metabolism³⁸. There is considerable evidence that the serotonergic neurotransmitter system is altered in depression³⁹ and ligands for measuring these receptors have recently been developed. A few studies have used PET to examine levels of serotonin receptors in depressed patients, but have reported conflicting results. In one experiment, 5-HT₂ binding was reduced in orbitofrontal areas in depressed patients⁴⁰, but no differences between depressed patients and controls were found in another study⁴¹. In older healthy adults 5-HT₂ receptor binding is reduced compared to young adults⁴², but is not further reduced in elderly depressed patients. However, serotonin binding is reduced in AD patients⁴³, indicating that results from studies of geriatric depression should be viewed with caution, as they may have been influenced by co-existing dementia, which is difficult to rule out clinically³⁹. It is clear that much work in this area remains to be done, including the assessment of brain function during serotonergic challenge in the elderly.

CONCLUSION

The accumulating evidence that detectable changes in brain function precede structural changes in neurodegenerative conditions and, in addition, that PET can provide a measure of severity of disordered brain function, indicates the potential value of PET in early detection and in monitoring the effects of treatments intended to modify the progression of these conditions. One important contribution of PET to the investigation of geriatric disorders will likely be in its use to delineate the mechanisms of disordered cerebral function. In addition, the use of pharmacological challenges and the measurement of neurotransmitter function hold promise for our understanding of neuropsychiatric disorders in the elderly.

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Single-photon Emission Computerized Tomography (SPECT)

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The introduction of single-photon emission computed tomography (SPECT) has markedly enhanced the study of brain function. The development of SPECT was the culmination of a series of investigations of cerebral blood flow (CBF) pioneered by Kety and Schmidt¹ in the late 1940s, combined with the introduction of transmission computed tomography (CT) in the early 1960s, in which three-dimensional images are derived from two-dimensional data. This chapter provides a review of the principles and basic techniques of SPECT, its present utility and application to clinical practice.

PRINCIPLES AND TECHNIQUES

Kety and Schmidt¹ pioneered CBF studies in man using nitrous oxide as a diffusible agent. These early studies required inhalation of nitrous oxide and sampling of arterial and internal jugular venous blood and could only provide a measure of whole-brain blood flow. Techniques to measure regional rCBF followed and used freely diffusible radionuclides, such as ⁸⁵krypton and ¹³³xenon, which were injected into the carotid artery². ¹³³Xenon emits γ -radiation, which can be detected through the intact skull, and multiple scintillation probes allow the measurement of CBF in specific regions of the brain. Further technological advances in detector sensitivity and data analysis allowed the replacement of intra-arterial injections by intravenous infusions or inhalation of gaseous ¹³³Xe, so that the measurement of rCBF became a non-invasive procedure. The rCBF data were initially presented in two dimensions and essentially reflected cortical flow, but with the development of the tomographic technology for SPECT, a three-dimensional measurement of rCBF could be obtained. The radiotracers used in SPECT emit a single γ -ray (or photon), as opposed to the dual simultaneous γ -rays of PET radiotracers, and hence the term "single-photon".

Recently, new radiotracers labelled with ¹²³iodine and ^{99m}technetium have been introduced. These lipophilic radiopharmaceuticals cross the blood-brain barrier and distribute in proportion to the rCBF shortly after intravenous injection. They are trapped in the brain and have a stable or static distribution over time, unlike freely diffusible, dynamic radiotracers such as ¹³³Xe, enabling images of higher resolution to be obtained. These agents can be used with the conventional rotating γ -cameras that are widely available in most nuclear medicine departments, whereas ¹³³Xe-labelled agents are not. The most advanced SPECT systems are now capable of a resolution of 8 mm and imaging times as short as 2–3 min for rCBF.

Unlike PET, there is no current prospect of SPECT being able to provide a direct measure of regional cerebral metabolism. However its use to measure neurotransmitter receptors is an evolving technique. The radioligand ¹²³I,3, quinuclidinyl 4-iodobenzilate (QNB) has been developed for the measurement of muscarinic acetylcholine receptors and has been applied to the study of Alzheimer's disease³.

NORMAL SUBJECTS AND NORMAL AGEING

Dynamic tomographic studies with ¹³³Xe produce similar values for grey matter flows as found with PET, but substantially overestimate white-matter rCBF and consequently give a limited distinction between grey and white matter. Static tomographic studies with ¹²³I- and ^{99m}Tc-labelled radiotracers cannot, as yet, provide an absolute measure of rCBF, but the pattern of rCBF, which can be expressed semiquantitatively, is similar to that found in PET studies, and the resolution is better than with ¹³³Xe dynamic tomographic studies.

The pattern of rCBF reported in normal subjects reflects, at least in part, the conditions of the subjects at the time. For example, if subjects are studied with eyes open, the visual cortex has the highest individual rCBF, but if the eyes are closed, the rCBF is reduced⁴. At present there are no generally accepted standard conditions for the control or resting state but it is clear that these should be standardized in any individual study.

CBF is reduced in relation to both advancing age and progressive cerebrovascular disease, and hypertension is the most important predisposing factor for a significant reduction in CBF.

The reduction in rCBF with age is confined to the grey matter—the white matter is unaffected and is more marked in anterior cortical regions. Females have higher rCBF than males for subjects up to 60 years of age, but the difference lessens above this age.

In the normal individual at rest, the rCBF is closely coupled to the regional metabolism of glucose and oxygen and is felt to reflect the underlying cerebral function. PET studies have shown that during visual stimulation, the rCBF and metabolic rate of glucose have similar marked focal increases in the visual cortex, while the regional cerebral metabolic rate of oxygen has a much smaller increase⁵. This uncoupling of glucose uptake and oxygen metabolism suggests that most of the extra glucose taken up during physiological stimulation is not oxidized and presumably lactate production by glycolysis increases. The implication of

these findings is that in normal resting individuals, rCBF and the regional cerebral metabolic rate of oxygen or glucose will provide similar information about the underlying cerebral function, but when physiologically (or perhaps pharmacologically) stimulated, they may convey information about different aspects of cerebral function, with none being an index of cerebral function. Similarly, in pathological cerebral conditions, close coupling of rCBF and the cerebral metabolic rates of glucose and oxygen cannot automatically be presumed to be retained.

STUDIES IN DEMENTIA

In addition to the effects of normal ageing, functional imaging studies in dementia have three further problems: (a) the normal variation in measures of cerebral blood flow or metabolism; (b) uncertainty of diagnosis of type of dementia (e.g. Alzheimer's disease (AD) or multi-infarct dementia (MID))—AD accounts for over 50% of cases, and can only be diagnosed definitively by cerebral biopsy or at autopsy; (c) the presence of cerebral atrophy. However, some demented individuals have little atrophy, while some normal subjects show considerable atrophy. Consequently, areas of apparently reduced rCBF in patients with dementia may be due to reduced flow to a normal volume of brain, normal flow

to a reduced volume of brain or reduced flow to a reduced volume of brain.

The studies by Kety and Schmidt¹ indicated lower rates of mean CBF in patients with dementia than in normal subjects, and most subsequent studies have confirmed these findings. Whole-brain studies have suggested that coupling between CBF and the cerebral metabolic rates of oxygen and glucose is present in the later stages of dementia, and although the link between CBF and the metabolic rate of oxygen is apparent in the early stages also, the metabolic rate of glucose is dissociated from these, such that it is relatively decreased in AD but increased in MID⁶.

The use of SPECT to study dementia has been reported since the mid-1980s. Using the ¹³³Xe inhalation technique, Bonte *et al.*⁷ found that out of 24 patients with probable AD, 19 had perfusion deficits, which were most commonly found in the parietotemporal flow, but instead had a patchy distribution pattern. However, ¹³³Xe SPECT involves measuring temporal variations in activity and is only possible with purpose-built equipment. The image characteristics are relatively poor because of scatter of the low-energy γ -rays and low counts obtained over the brief scanning periods.

The radiopharmaceutical ¹²³I-iodoamphetamine (IMP) does not present these technical problems and its initial cerebral uptake is proportional to rCBF. It was the first tracer to be

Table 67.1 ^{99m}Tc-HMPAO SPECT studies in dementia

Study	Patients (n)	Normal controls	Diagnosis	Assessment of images	Results
Neary <i>et al.</i> (1987) ¹¹	AD (23) FLD (9) PSP (9)	None	Clinical, NINCDS-ADRDA, CT scan	Qualitative	AD: posterior Cr Cerebral abnormalities, FLD and PSP anterior Cr abnormalities
Perani <i>et al.</i> (1988) ²⁵	AD (16) early	16	NINCDS-ADRDA, CT scan/MRI neuropsychology	Semiquantitative ratio of ROI to cerebellum	Decreased rCBF in frontal and temporo-parietal areas, hemisphere asymmetry
Burns <i>et al.</i> (1989) ²⁶	AD (20)	6	NINCDS-ADRDA, CT scan, psychometry	Semiquantitative ratio of ROI to cerebellum	Decreased rCBF in temporal and posterior parietal areas, regional rCBF correlations with memory, praxis and language functions
Gemmell <i>et al.</i> (1987) ¹⁰	AD (17) MID (10)	3	DSM-III, Hachinski, MRI	Qualitative	Perfusion defects (temporoparietal occipital) more common in AD than MID, more bilateral defects in AD than MID
Battistin <i>et al.</i> (1989) ²⁷	AD (21) Mixed (9)	None	DSM-III, CT Scan	Semiquantitative ratio of ROI to cerebellum	Decreased rCBF (parietal) in AD than mixed AD and MID form, few showed right-left asymmetry
Goldenberg <i>et al.</i> (1989) ¹⁷	AD (23)	None	Clinical CT scan neuropsychology tests	Semiquantitative ratio of ROI to all ROIs	Co-variation of neuropsychology test results with frontal inferior parietal and superior temporal regions
Podreka <i>et al.</i> (1987) ²⁸	Dementia (12)	None	CT scan	Semiquantitative (region/whole brain)	All patients showed perfusion defects
Upadhyaya <i>et al.</i> (1991) ²⁴	AD (15)	10	NINCDS-ADRDA Geriatric Mental State, CAMCOG	Qualitative, semi-quantitative (ROI/cerebellar)	All AD patients showed perfusion defects Significantly low ratios in parietal, frontal and occipital areas.
Jobst <i>et al.</i> (1998)	Dementia (200) Annual evaluation to necropsy	119	NINCDS-ADRDA DSM-III-R	Semiquantitative	AD had medial temporal lobe atrophy (CT Scan) and parietotemporal hypo-perfusion. Both markers had better diagnostic accuracy than NINCDS-ADRDA and DSM-III-R.
Shih <i>et al.</i> (1999)	AD (18) Repeated 5-23 months	None	MMSE	Semiquantitative	All AD had decreased CBF in consecutive SPECT
Defebvre <i>et al.</i> (1999)	DLB (20) AD (20) PD (20)	None	MMSE	Semiquantitative	DLB: frontal CBF lower than AD
Mullet <i>et al.</i> (1999)	AD (116)	20	DSM-III-R MMSE	Semiquantitative	3 regions with decreased CBF AD: 48% Controls: 10%

widely used in static SPECT studies of dementia. These have generally reported that patients with AD have deficits in flow which are maximal bilaterally in the parietotemporal cortex, while patients with MID vary from having a normal pattern to marked asymmetrical focal deficits anywhere in the cortex. Sharp *et al.*⁸ found that, despite having characteristic regional abnormalities on SPECT, the majority of patients with AD had a normal appearance in those regions on magnetic resonance imaging (MRI). Ebmeier *et al.*⁹ found that while MRI was not able to differentiate between AD and MID, 19 out of 21 patients showed bilateral deficits on IMP SPECT scans. Although the distribution of ¹²³I-IMP initially reflects the rCBF, redistribution of the tracer occurs approximately 1 h after injection. The use of ¹²³I-IMP is also limited by the restricted availability of ¹²³I, which has a half-life of 13 h, and is costly to produce, as this requires a cyclotron.

^{99m}Tc-HMPAO SPECT

^{99m}Technetium is readily available from commercial generators in nuclear medicine departments, is inexpensive and has a shorter half-life (6 h) and better dosimetry than ¹²³I. Labelling with ^{99m}Tc is a much more complex procedure than with ¹²³I and the development of a ^{99m}Tc-labelled compound to provide a measure of rCBF was a major advance in SPECT technology. Technetium-labelled hexamethyl propyleneamine oxime (HMPAO) is a lipophilic tracer which, after intravenous administration, crosses the blood-brain barrier with high extraction and is retained in the brain in hydrophilic form. The brain uptake occurs over the first 2 min and has a stable distribution for many hours. This enables conventional equipment to be used to detect the radiation emitted from the brain. Comparative studies with PET in humans reveal a tendency to underestimate flow in areas of high rCBF but a good correlation with areas of low and medium rCBF. Although ^{99m}Tc-HMPAO cannot, at present, be used to quantify rCBF absolutely, the results can be expressed semi-quantitatively by comparing the counts in each brain region of interest (ROI) to a reference area, such as the whole brain or cerebellum.

A number of SPECT studies have now been performed using ^{99m}Tc-HMPAO to investigate dementia and the characteristics of these are shown in Table 67.1. The studies varied in the populations studied, diagnostic criteria used and the methods of assessment of images. Some of these studies had no data on control subjects for comparison. The majority carried out initial screening with CT or MRI to exclude structural lesions.

The early studies used qualitative assessment of the images and found that most patients with AD had bilateral parietotemporal deficits; those who did not tended to be less impaired cognitively¹⁰.

Frontal deficits were also seen in AD. Deficits were less common in MID than AD, particularly when known infarcts on MRI were excluded, and they occurred in a more variable and asymmetrical pattern. Neary *et al.*¹¹ also investigated patients with progressive supranuclear palsy (PSP) and others with the clinical syndrome of dementia of frontal-lobe type, which is consistent with, although not necessarily diagnostic of, Pick's disease.

The later studies have generally been semi-quantitative. When a rotating γ -camera has been used, the whole brain is scanned and the cerebellum has usually been chosen as the reference area, as it is relatively unaffected by the pathology of AD¹². Also, patients with AD have a normal cerebellar metabolic rate of glucose, compared with elderly controls, on PET study¹³. A recent study validated the use of the cerebellum as a reference region for SPECT quantification in patients with AD¹⁴. Studies using dedicated head scanners, which acquire their data in the form of transverse tomographic slices, have not usually obtained cerebellar data, so they have used the occipital region as a reference area because the latter also appears relatively unaffected by AD¹⁵.

These studies have consistently found that patients with AD have a bilateral decrease in rCBF in the posterior temporal and parietal regions, adjacent to the occipital lobe, and sometimes the frontal lobes are affected, although the basal ganglia are not (Table 67.1). Several studies have demonstrated correlations in AD between rCBF and neuropsychological function, and these have tended to be more apparent for language, praxis and global function than for memory^{15,19}. Recent studies of AD have also evaluated the correlates of changes in CBF in relation to psychopathology and behaviour disturbance. Reduced frontal CBF in AD is associated with negative symptoms²⁰, with presence of delusions²⁴, disinhibited behaviour, apathy and blunted affect²².

Pharmacological activation studies of the effect of central cholinergic stimulation upon rCBF in AD have recently been reported. These require paired SPECT studies, basal (after saline infusion) and activated (after infusion of a cholinergic agent, e.g. physostigmine). In patients with AD, the reduced rCBF in the posterior parietotemporal region in the basal scan was focally increased in the scan after the cholinergic agent, which did not occur in control subjects¹⁹. AD is associated with striking abnormalities in the central cholinergic system, of a functional cholinergic deficit that is at least partly reversible. Also a SPECT study of nootropic drugs showed regional improvement in CBF²³. Donepezil treatment in AD was associated with a significant increase in CBF in the frontal lobes²⁴.

Studies have also addressed the diagnostic value of SPECT in differentiating AD from other types of dementia. Dementia with Lewy bodies (DLB) is associated with reduced frontal CBF than AD²⁵. Importantly DLB is associated with severe degeneration of the dopamine system as shown by a recent SPECT study²⁶. Primary progressive aphasia is associated with reduced CBF in fronto-temporal regions²⁷. Fronto-temporal dementia FTD is characteristically associated with reduced CBF in fronto-temporal regions²⁸. Alzheimer's disease and FTD could be differentiated by discriminant analysis applied to ^{99m}Tc-HMPAO SPECT data with 100% correct classification of patients with FTD and 90% correct clarification of patients with AD²⁹.

Moreover fronto-temporal dementia is characterized by early non-cognitive behavioural changes with relatively spared cognitive function, frontal atrophy and SPECT deficits in CBF in frontal and temporal regions whilst AD is associated with early cognitive changes and medial temporal and parietal-temporal deficits in CBF³⁸.

Patients with Korsakoff's psychosis, in contrast to patients with AD, have normal rCBF in the posterior temporal region.

It is well established that Parkinson's disease (PD) is characterized by degeneration of dopaminergic neurons in the substantia nigra, which project to the corpus striatum. In an HMPAO SPECT study, 36 patients with PD were found, overall, to have normal rCBF in the caudate and putamen; however, those on no therapy had lower rCBF, and those on L-dopa-replacement therapy had higher rCBF, in the caudate and putamen, than controls. Bilateral reductions in rCBF in the parietal region, similar to those found in AD, were also found in patients with PD. No patients had focal lesions on CT scans. Interestingly, the parietal reductions in rCBF were more pronounced in those PD patients with cognitive impairment and those receiving chronic anticholinergic therapy. Dopamine deficiency is the main, but not the only, neurochemical deficit in PD, and these SPECT findings are consistent with the concept that AD and PD may overlap to some extent. Patients with Huntington's Disease (HD), characterized by chorea and progressive dementia, have reduced rCBF in the caudate nucleus, matched by caudate atrophy on MRI. However, the size of the rCBF deficit in the SPECT study exceeded that predicted by the MRI findings³⁹.

There has been a case report of a patient with pathologically proven Creutzfeldt-Jakob disease, who had strikingly reduced

rCBF throughout the cortex on HMPAO SPECT study, but a normal CT scan⁴⁰. Patients with the AIDS dementia complex have been found, using SPECT, to have multiple or focal rCBF deficits, correlating with focal signs or symptoms, while CT scans showed diffuse cerebral atrophy⁴¹.

DISCUSSION

Studies employing SPECT and PET can reveal cerebral abnormalities when CT and MRI do not, because the latter are measures of cerebral structure, while the former are measures of cerebral function. SPECT is, and is likely to remain, much more generally available than PET, but it is not currently capable of absolute quantification.

The characteristic SPECT findings in AD are bilaterally decreased rCBF in the parietal and temporal lobes adjacent to the occipital lobes, sometimes involving the frontal lobes, particularly in later cases. The primary motor, sensory and visual cortices and basal ganglia are relatively unaffected. This contrasts with typical CT findings of diffuse cerebral atrophy, although the CT scans have not generally been orientated along an axis that obtains optimum views of any focal atrophy in the hippocampus and temporal lobe⁴². However the Oxford Project to Investigate Memory and Aging (OPTIMA), which involved 200 patients with dementia and 119 normal controls evaluated annually till necropsy showed that medial temporal lobe atrophy (CT Scan), (80% diagnostic accuracy) and parietotemporal hypoperfusion by SPECT (83%) predicted the pathology of AD better than the established clinical criteria of NINCDS-ADRDA (66%) and DSM-III-R (66%)³⁵. On careful assessment, focal features can be seen on CT scans which correlate with individual symptoms, such as aggression or wandering⁴³, but the extent of the focal CT abnormalities is less than that seen on SPECT, and it appears that structural atrophy lags behind clinical deficit.

The diagnosis of dementia and its differentiation from other clinical syndromes, such as depression, is still primarily a clinical matter. If a structural lesion, such as a cerebral tumour, is suspected, then a structural scan, such as CT, is likely to be the best test for this. What, then, is the appropriate role of SPECT in current clinical practice in the assessment of dementia? The study by Upadhyaya *et al.*³⁴ suggests that it would be unwise to expect SPECT to assist in the differentiation of dementia from depression in the elderly because similar, albeit less marked, changes are seen in depression, as in AD. It would be interesting to know if those elderly individuals who present with the clinical syndrome of depression, and who have the characteristic AD abnormalities on SPECT, have a worse prognosis than those with normal SPECT scans. They might be individuals at risk of developing AD where depression has been one of the early clinical features, but much more information, including follow-up studies, is needed on this patient group to evaluate the utility of SPECT. More information is also needed on the frequency with which perfusion deficits (particularly those with a pattern similar to that in AD) are seen on visual inspection of SPECT studies in healthy, elderly controls. Patients with PD, particularly those who are cognitively impaired or receiving chronic anticholinergic treatment, also have similar SPECT abnormalities to patients with AD. Perhaps it is not surprising that there may be similarities on this measure of brain function, as there can be a degree of overlap of the clinical syndromes.

If the clinical diagnosis of dementia is made in an individual patient and it appears to be a primary degenerative dementia, the SPECT study can add weight to the clinical impression that the underlying diagnosis is that of AD, frontal lobe dementia (FLD) or progressive supranuclear palsy (PSP). AD is most characteristically associated with posterior rCBF deficits, while FLD and PSP are strongly associated with anterior rCBF deficits and

themselves have distinctive patterns when CT scans are not particularly helpful in this situation. Similarly, Korsakoff's psychosis appears to be more characteristically associated with frontal rCBF deficits than posterior temporal deficits¹⁷. SPECT can also assist the clinical assessment in the differentiation of AD from MID: bilateral parietotemporal deficits are strongly suggestive (but not diagnostic) of AD, although it must be remembered that AD and MID coexist in a substantial minority of cases. There is no particular single pattern of rCBF deficits seen in MID. SPECT findings vary from normal to asymmetrical deficits and theoretically MID could mimic any other pattern. However, if an asymmetrical pattern is found affecting areas other than the parietotemporal region, it is suggestive of MID. This impression is strengthened if the rCBF deficits coincide with cerebral infarcts seen on structural imaging with CT or MRI. If a patient with relatively advanced dementia has a normal SPECT, it is unlikely to be AD and may be consistent with MID.

All these conclusions, however, must be tempered by the knowledge that our present association of SPECT and PET patterns with specific conditions is based on studies where clinical diagnoses of patients are made initially and are then correlated with the SPECT findings. We need follow-up studies and pathological findings at post mortem to make more definitive assessments of the significance of specific SPECT patterns. Indeed a recent study evaluated early AD (mild cognitive impairment) using brain perfusion SPECT who were diagnosed to have AD 2 years later with a follow-up SPECT⁴⁴. Selective reduction in CBF was observed in the left hippocampus and parahippocampal gyrus in the follow-up SPECT. We could also learn whether SPECT studies can help predict clinical outcome in individual cases. There is evidence from PET studies of patients with early AD that focal cortical reductions in glucose metabolism precede the appearance of focal neuropsychological deficits.

However, if the early whole-brain studies are correct and the metabolic rate of glucose is uncoupled from CBF in early AD, such that it is decreased relative to CBF and the rCBF is reduced later in the course of the disease, SPECT might have different predictive power in comparison with PET in early AD. It is possible that patients at risk for HD may be diagnosed in the presymptomatic phase by the SPECT finding of reduced rCBF in the caudate nucleus.

It is clear that SPECT will be used substantially in further research in AD. More significant advances in our knowledge of AD are likely to come from follow-up and activation studies. The latter would involve paired SPECT studies, basal and activated, and the activation could be cognitive or pharmacological. For paired studies, the important measure is the difference in rCBF between the two SPECT studies, which indicates the effect of activation. This avoids the problem of variation in basal values between individuals and the problem of the effect of an uncertain degree of cerebral atrophy upon rCBF. The semi-quantitative measures of SPECT are adequate, and may be preferable, for this analysis. Cognitive activation studies in controls and in patients with AD would provide a better understanding of the cerebral processes involved in cognitive function in normal individuals, and knowledge about the form of disruption of these processes in disease. Pharmacological activation studies would enable direct measures of specific neurotransmitter function to be made and hence the pharmacological characterization of disease in an individual, with implications for treatment and prognosis¹⁹.

Another promising direction for SPECT is the study of neurotransmitter receptors. The radioligand ¹²³I-QNB has been developed for the measurement of muscarinic cholinergic receptors and it has been used in the study of AD³. Currently it requires a radiochemist on site to produce ¹²³I-QNB, and so it is not commercially available, but the development of radioligands for

general use in receptor studies is likely to continue and may further extend the role of SPECT.

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Part F

Affective Disorders

- FI Nosology and Classification
- FII Depression, Dysthymia, Bereavement
and Suicidal Behaviour
- FIII Mania

Nosology and Classification of Mood Disorders

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The mood or affective disorders are a group of disorders characterized by disturbance of mood and accompanied by a partial or complete change in mood or affect that is either manic (or hypomanic) or depressed (or mildly dysphoric). These disorders are classified in both the DSM diagnostic system and the ICD diagnostic system in the three-digit rubric of 296^{1,2}. There have been recommendations for changes in the ICD system from ICD-9 to ICD-10 that would reclassify some disorders previously classified elsewhere (such as neurotic depression, 300.4) to the mood or affective disorders. Another minor change in the new ICD diagnostic system (already represented in DSM-IV) is replacing the title “affective disorder” (affective psychosis) with “mood disorder”. With some exceptions noted below, the nosology of mood disorders in ICD-10 and DSM-IV is appropriate for the classification of mood disturbances in older adults. The classification of the disorders will therefore be described below, with specific comments related to older adults.

MANIC EPISODE

A manic episode is characterized by an elevated mood that is unrelated to the patient's circumstances. This elevated mood usually varies from an expansive or irritable syndrome to an almost uncontrollable excitement and psychotic agitation. Changes in mood are accompanied by increased energy, a decreased need for sleep, a decline in normal social inhibitions, as well as inflated and grandiose ideas, which frequently become delusional. Older persons can experience a typical episode of mania as well as hypomania (a less severe elevation of mood) but are more likely to suffer a so-called “irritable” or “angry” manic episode. Joviality and elation are replaced by irritability and agitation (as described elsewhere in this text). Nevertheless, older persons who suffer manic episodes usually meet both ICD-10 and DSM-IV criteria, even when the predominant symptoms are irritability and anger, as other diagnostic criteria are met.

DEPRESSIVE (OR MAJOR DEPRESSIVE) EPISODE

A major depressive episode is characterized by a depressed, disinterested or irritable mood, associated with the loss of interest or pleasure in all or almost all activities, accompanied by a number of other symptoms. These additional symptoms include a reduced capacity for enjoyment, reduced interest in surroundings, difficulty concentrating, lethargy, sleep distur-

bances, appetite disturbances, decreased self-esteem and self-confidence, and frequent ideas of guilt or worthlessness. Although older persons are somewhat less likely to report a specific decrease in their mood, they almost always describe a loss of interest in usual activities (anhedonia) in the midst of a major depressive episode. The categories of severe depressive episode in ICD-10 and major depressive episode in DSM-IV are very similar. DSM-IV criteria permit the diagnosis of a minor depressive episode, that is, a depressive episode which fulfills some of the symptom criteria for a major depressive episode and/or dysthymia and which lasts 2 weeks or longer. The more severe consequences of a depressive episode, such as a successful suicide or a retardation that progresses to stupor, would be characteristic of a severe but not a mild depressive episode in ICD-10.

Older persons who suffer a complicated, severe depression are easily diagnosed according to both ICD-10 and DSM-IV criteria. Problems do arise, however, when the episode experienced by older persons is accompanied by a severe medical illness or significant cognitive impairment. The frequency of psychobiologic symptoms in more severe depressions renders the distinction between symptoms of depression and symptoms of physical illness/functional impairment difficult in the midst of a depressed mood associated with medical illness. The current nosology is not helpful in disaggregating mood disorders from either the symptoms of physical illness or normal psychologic reactions to physical illness. Some have suggested that a unique rating for older persons for depression should include a measure of cognitive functioning (but this is yet to be included in any extant diagnostic system). The co-morbidity of depressive symptoms in organic mental disorders such as dementia, Parkinson's disease and stroke renders such an approach potentially useful in improving the current nosology. Major (or severe) depression can be specified as severe, with or without psychotic features in DSM-IV. Psychotic depression is relatively more frequent in the elderly and the recognition of psychotic features may direct the specific therapeutic intervention considered by the clinician.

BIPOLAR AND UNIPOLAR DISORDER

In both ICD-10 and DSM-IV, individuals who present with a history of recurrent mood disorder (at least two), in which mood and activity were profoundly disturbed and at least one of the episodes is manic, are classified as suffering from bipolar mood disorder. Bipolar disorder usually has an early age of onset, although it can have its onset later in life, and the categorization is useful in older as well as younger persons. Bipolar disorders are

classified as manic (the individual is currently in a manic episode), depressed (the individual is currently in a depressed episode and has at least one well-authenticated manic episode in the past), or mixed (the individual's current episode involves a full symptomatic picture of both manic and depressed episode, except for the duration requirement of 2 weeks of depressed episodes). Symptoms are often intermixed or rapidly alternating within hours or days in the latter category.

Bipolar II disorder is characterized by at least one hypomanic episode and at least one episode of major depression. These disorders are probably more common among older persons than the more classic bipolar disorder described above. Older persons frequently experience depression and occasionally mania secondary to medical illness or medications, categories which are specified in DSM-IV as mood disorder due to a general medical disorder and substance-induced mood disorder. Common examples of causes of these secondary mood disorders in the elderly include hypothyroidism, Parkinson's disease, chronic obstructive pulmonary disease, various forms of cancer, alcohol, β -adrenergic blockers and L-dopa. Mood disorders with a seasonal pattern are less frequent in the elderly.

Individuals who never experience an episode of mania or hypomania but nevertheless suffer episodes of major depression are generally classified as "single-episode" or "recurrent". Most depressive disorders are recurrent, that is, if an individual (regardless of age) suffers an episode of major depression, the chances are high that he/she will suffer an additional episode at some time in his/her life. Recurrent or persistent depressive episodes are not always easily classified, that is, distinct episodes of major depression followed by distinct episodes of normal functioning may be the exception rather than the rule. For this reason, additional diagnostic categories have been instituted in the nosology to disaggregate the classification of the natural history of depressive episodes. In ICD-10, a category of "recurrent severe depressive disorder" is analogous to the DSM-IV category of "major depression, recurrent". Recurrent mild depressive disorder is categorized in ICD-10 by repeated episodes of depression, the majority of which fulfill the description given for mild depressive episode above. These recurrent mild episodes may or may not be associated with environmental stress and may be either acute or insidious in onset. Recovery is usual, although not always complete.

Cyclothymia is included in both DSM-IV and ICD-10. According to ICD-10, cyclothymia is a persistent instability of mood involving numerous periods of mild depression and mild elation. The instability usually develops early in adult life and pursues a chronic course, although at times mood may be normal and stable for months. Cyclothymia is difficult to establish without a prolonged period of observation or an unusually good account of the subject's past behavior. In DSM-IV, cyclothymia must persist for at least 2 years to be diagnosed and involve numerous hypomanic episodes.

Yet another phenomenon has emerged in the study of the natural history of depression—rapid cycling. Rapid cycling is the occurrence of four or more episodes of depression or mania per year, with either 2 weeks of normal mood between episodes, or a shift directly from mania to depression, or vice versa. Others have classified rapid cycling as involving either mixed mood states, or frequent mood fluctuation without discrete intermorbid periods, or involving 24 or 48-h cycles of mood disturbance. Rapid cycling occurs in persons regardless of age, although there may be some tendency for individuals to decrease their propensity to rapid cycling with increased age. Rapid cycling appears to be more common in individuals with bipolar disorder than unipolar disorder, but it is not limited to individuals with bipolar disorder. Rapid cycling is a specifier of bipolar mood disorder in DSM-IV.

DYSTHYMIA

Dysthymia is a category included in both ICD-10 and DSM-IV. Dysthymia is a chronic depression of mood which does not fulfill the description and guidelines of mild recurrent depressive disorders (in ICD-10) and is characterized by periods of days or weeks at a time when patients feel tired and depressed, where everything is an effort and where nothing is enjoyed. They "brood and complain", sleep badly and feel inadequate, but are usually able to cope with the basic demands of everyday life. This description is similar to the description of dysthymia in DSM-IV, where the essential feature is a chronic disturbance of mood, involving a depressed mood for most of the day and more days than not for at least 2 years (the 2 year minimum duration is not a criterion in ICD-10). In addition, individuals suffering dysthymia suffer many of the symptoms of the more severe depressions, but with less severity, such as poor appetite, insomnia, low energy and low self-esteem. Older persons often meet criteria for a dysthymic disorder when diagnosed with major depression. It is often difficult to determine, however, whether the dysthymic disorder is a separate entity or whether it is part of the same syndrome which "waxes and wanes" in severity. Recent studies have suggested that dysthymic disorder may exist most often in the elderly as an entity not associated with major depression. Dysthymic disorder has usually been equated with depressive neuroses, as described in earlier versions of both the ICD and DSM diagnostic systems, which assumes that the depressed mood results from psychoneurotic and stress-related difficulties.

Yet another category which has been suggested is minor or mild depression. Criteria for minor depression can be found in the appendix of DSM-IV and variants can be found in the Research Diagnostic Criteria and in ICD-10. There is virtually no data to substantiate a specific entity for minor depressions, except that persons who experience minor depression (that is, symptoms less severe than major depression for at least 2 weeks, according to DSM-IV) exhibit a similar risk factor profile and are at much greater risk for developing major depression over time. There have been a number of studies of minor depression (see Chapter 71) among the elderly. The renewed interest in minor depression once again brings the categorical vs. continuum controversy to the forefront in the phenomenology of depression.

ADJUSTMENT DISORDER WITH DEPRESSED MOOD

In the DSM-IV system of classification, an adjustment disorder is a reaction to an identifiable psychosocial stressor that leads to maladaptive reactions, including impairment in occupational functioning and symptoms that are in excess of normal and expected reaction to this stressor. The maladaptive reaction may take many forms and an exaggerated depressed mood is one of the forms.

Symptoms of adjustment disorder with depressed mood include tearfulness and feelings of hopelessness. By definition, the maladaptive reaction can persist no longer than 6 months, does not meet criteria for other disorders and does not represent uncomplicated bereavement. ICD-10 includes the category of adjustment disorder and the clinical form may take the characteristic of a "prolonged depressive reaction", that is, a mild depressive state occurring in response to prolonged exposure to a stressful situation but of a duration not lasting 6 months. The categories of adjustment disorder are therefore very similar for both diagnostic systems. In theory, the stressor may include physical illness and therefore this category is most relevant to depressive episodes in later life. A lack of acceptable longitudinal data on the association of depressive symptoms with physical

illness renders it difficult to determine whether older persons do recover from the depressive episode, which often accompanies a physical illness within the 6 months required by the diagnostic category of adjustment disorder.

DEPRESSED MOOD ASSOCIATED WITH DEMENTIA OF THE ALZHEIMER'S TYPE AND VASCULAR DEMENTIA

DSM-IV provides for the classification of dementia of the Alzheimer's type (both early and late onset) as "with depressed mood as well as vascular dementias". The frequency of co-morbid depression and cognitive impairment renders this classification a common one among dementia patients. As the actual pathophysiological and psychopathological relations between depression and cognitive impairment are unclear, the simple recognition of the co-morbidity is sufficient. For example, the depression may be a reaction to cognitive decline or an actual symptom of the underlying disease process, which also causes the cognitive problems. Some recent studies have suggested that a separate category of vascular depression be introduced. This condition is present when individuals meet the criteria for major depression and have MRI-confirmed vascular brain changes. Persons with vascular depression appear to have more symptoms of apathy (perhaps due to disruption of prefrontal systems or their modulating pathways) and may be at increased risk of non-recovery.

CONCLUSION

In general, the current classificatory system of both DSM-IV and ICD-10 work relatively well for classifying older persons suffering from mood disorders. A number of distinct exceptions must be recognized, however. Categories such as dysthymic disorder, adjustment disorder and minor depression in DSM-IV do not appear to be adequate, and therefore more exploration of the ICD-10 construct of mild depression (but possibly not utilizing the specific diagnostic criteria of ICD-10) would appear in order. A major problem with the current classification systems is the inability to take into account co-morbidity, especially co-morbid

depression and physical illness. Co-morbid depression and physical illness is a grossly unstudied area, compared to the clinical relevance of the condition. To what extent do our current classification systems accommodate individuals suffering mild or even severe depressive symptoms in the midst of physical illness? In addition, depression is often co-morbid with other psychiatric symptoms, especially anxiety disorders and somatic complaints. Neither ICD-10 nor DSM-IV adequately accommodates the co-morbid psychiatric syndromes that are frequently seen in older adults.

Finally, when classifying mood disorders, many individuals suffer a depressed mood in late life that is not disordered. Uncomplicated bereavement is an expected accompaniment of older persons who experience a significant loss in old age. In addition, other older persons may become demoralized, given the current circumstances in their lives. Such persons should not be classified in a disease-orientated classification system. Nevertheless, these human experiences are not to be ignored by the clinician working with the older adult suffering a mood disorder.

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Genetics of Affective Disorders

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The observation that affective disorders aggregated in families has been present from the time of Socrates, through the analysis of Sigmund Freud, the behavioral work of B. F. Skinner and in the genetic studies of the past 40 years. In each of these views, the family unit recurs as a major focal point in mood disorders. But how is one to understand this relationship? Affective disorders may be aggregated in families for a variety of reasons, such as shared genes, shared culture, shared adversity, or multiple other reasons.

In the past, research in families followed two non-overlapping tracks: genes and environment. This was known popularly as “nature vs. nurture.” In recent years, researchers have become more aware that attempts to dichotomize the cause of affective disorders into “genes” and “environment” has hindered understanding of mood disorders. The emerging science of genetic epidemiology focuses on causes, distribution and control of disease in groups of relatives, and with inherited causes of disease in populations¹. Thus, the goal of genetic research is to identify both genetic and environmental causes of illness. In fact, certain types of genetic studies (e.g. twin studies) have provided some of the strongest evidence that environment plays a significant etiologic role in the expression of mental illness.

Research findings clearly show that the etiology of mood disorders in early and middle adulthood is multifactorial. Both environmental factors (e.g. parental loss, stroke) and genetic factors are strongly implicated, although what occurs and how this happens remain as areas of intense research. In recent years, investigations in late-life affective disorders have asked the same questions regarding cause, but the answers suggest that there are etiologic differences among affective disorders at different ages. This observation only leads to new questions. Is the development of late-life affective disorders related to the genetic make-up of the individual? Are late-life affective disorders a continuation of mood disorders which developed earlier in life or do they represent unique reactions to specific challenges presented by the process of aging? If so, what are these challenges and how are they related to an individual’s genetic make-up? How do genetic and environment factors contribute to the understanding of diagnostic subsets of affective disorders?

In this chapter, we will review the current state of genetic research of affective disorders, primarily studied in early and middle life. Then we will contrast that with what is known about affective disorders that have their first onset in late life. Finally, we will discuss the implications of these findings and suggest a theoretic framework to understand genetic and environmental contributions to geriatric affective disorders.

GENETIC INFLUENCES IN AFFECTIVE ILLNESS

There are four types of studies that demonstrate the influence of genetic factors on the development of disease: (a) studies of familial aggregation; (b) twin studies; (c) adoption studies; and (d) association and linkage studies of an illness with a genetic marker.

Family Studies

Family studies are the most basic research form in evaluating genetic risk. They attempt to answer the question, “Does this disorder run in families?” Logically, if genes cause a disorder, then relatives of an individual affected by that disorder (the proband) should be more likely to have that particular disorder than others in the general population, since they share common genes. However, common environmental factors may also contribute to the clustering in families. Therefore, the major goal of family studies is to understand the magnitude and patterns of a disease’s aggregation, rather than identify specific genetic causes.

In affective disorders, six landmark studies²⁻⁷ have supported the observation of familial aggregation. First-degree relatives (parents, siblings, offspring) of depressed patients are twice as likely to develop depression than those in the general population. First-degree relatives of bipolar patients are at even greater risk for developing bipolar disorder (incidence of 3.7–17.5%, depending on the study). The risk of bipolar disorder is less in second-degree relatives (grandparents, grandchildren, aunts, uncles) of bipolar patients, although still elevated over the general population.

Twin Studies

Twin and adoption studies are especially helpful in answering the next question, “What are the relative contributions of genes and environment to the development of a mental disorder?” Twins provide a natural experiment in genetics. Monozygotic (MZ) twins, known as identical twins, share 100% of their genes. Dizygotic (DZ) twins, known as fraternal twins, share approximately 50% of their genes, the same proportion as with other siblings. In the “twin experiment”, it is assumed that both dizygotic and monozygotic twins share the same environment, but only monozygotic twins share the same genes. Thus, if genes are a putative cause for a disorder, then the MZ co-twin of a proband should be at higher risk to develop a disorder than the DZ co-twin of a proband. This is expressed as the concordance rate.

In affective disorders, the average concordance rate for bipolar disorders in monozygotic twins was 60%, while the concordance rate in dizygotic twins was 12%⁸. The five-fold greater rate of concordance strongly indicates the importance of genetic factors in the familial aggregation. For major depressive disorders, the role of genes is much weaker. Two recent twin studies^{9,10} found that the relative risk for monozygotic twins was 1.9, while the risk for dizygotic twins was 1.2. Both findings of heritability were statistically significant.

Adoption Studies

Adoption studies provide a powerful method of evaluating genetic and environmental contributions to the familial aggregation of disease. In these cases, adopted children have a genetic relationship with their biological family, but share the primary environmental relationship with their adopted family. Thus, if genes are responsible for the familial transmission of a disorder, then the risk for the adopted child should more closely match the risk of the biological family. However, if environmental factors are more responsible, the relative risk of the adopted child would match the risk of the adoptive family.

In affective disorders, five studies have reviewed adoption and incidence of affective illnesses¹¹. Two of these studies have demonstrated strong genetic influences, while the others show a more variable response. One adoption study¹² suggested certain environmental influences, especially parental loss and parental alcoholism, as predictors of depression.

Genetic Markers and Linkage Studies

The most specific studies for genetic involvement are genetic marker and linkage studies. Genetic markers are specific areas on the chromosome where laboratory procedures may differentiate individuals on the composition of DNA at that location. They are used to identify where a specific gene may be on a chromosome. Linkage refers to the nearness of two or more genes or markers on a chromosome. Linkage studies are based on the principle that as the genetic distance decreases, the probability that they will be inherited increases.

A number of gene loci for affective illness have been proposed through linkage work, but the results have been disappointingly inconsistent, and a number of the most striking findings have proven controversial¹³. One possible exception is a small subset of bipolar disorders that may be X-linked^{11,14}. Recent investigations in an ethnically homogeneous American Amish population initially suggested linkage to a marker on chromosome 11¹⁵. Unfortunately, more recent work has cast doubt on this conclusion¹⁶ and has prompted greater caution generally in the interpretation of the linkage results in psychiatry.

LATE-LIFE AFFECTIVE DISORDERS

Just as affective disorders in early and middle life are quite heterogeneous, late-life mood disorders are also a heterogeneous entity. However, it is unclear whether late-life affective disorders comprise the same notions of heterogeneity as those in early and middle life. In late-life mood disorders, two groups of patients are usually differentiated: those who had an early-onset of illness and continue to have symptoms or episodes in their later life, and those who had a late-onset of illness (usually defined as a first onset after the age of 60 years).

Mendlewicz¹⁷ and other researchers have suggested that the age of onset is an important clinical variable in the genetic study of

affective illness. When differentiated by age of onset, late-onset depressive disorders have qualities that suggest they may be genetically different from early-onset forms. These characteristics include the non-conformity of affective disorders to expected genetic models and clinical differences observed between late- and early-onset depression.

Most genetic models propose that strong genetic influences are most often found in the most severe forms of an illness and in the early expressions of the illness. However, late-onset affective illness syndromes are typically among the severest and most refractory to treatment^{18,19}. Older patients with affective disorders who require hospitalization tend to have a slower resolution of symptoms and a longer duration of hospitalization than younger adult patients²⁰. Further, patients with late-onset depression and bipolar disorders have less familial aggregation of mood disorders than those with the early-onset affective disorders. This finding has been extensively validated in many studies²¹⁻²⁶. Late-onset mood disorders are also less likely to display the gender disparity²¹ present in early-onset mood disorders.

Clinically, patients with late-onset mood disorders tend to have increased impairment on neuropsychological testing and higher rates of incident dementia noted at follow-up²⁷. While inadequate support appears particularly difficult among middle-aged patients, stressful life events appear to pose the greater risk of poor outcomes among geriatric patients²⁸.

Gatz *et al.*²⁹ studied the characteristics of depression in a sample of elderly reared-apart and reared-together Swedish twins. They found evidence of only limited heritability (16%). In contrast a study³⁰ of a sample of Danish twins 75 years of age and older found that depression symptomatology is moderately heritable in late life (approximately 35%). However, both studies consistently implicated environmental factors as the major source of variance in depression symptoms among the elderly.

GENETICS OF LATE-LIFE AFFECTIVE DISORDERS

How can one understand the differences noted above, and how are they important for the genetic study of late-life mood disorders? Two explanations have been proposed for the age-related variation in genetic influence for affective illnesses: multifactorial inheritance and increased phenocopy expression with age. Multifactorial inheritance (also called polygenic inheritance) suggests that inheritance patterns are a result of a combination between multiple interacting genes and environmental factors. Examples of traits that have multifactorial inheritance include stature, intelligence and blood pressure¹⁷.

According to the multifactorial theory, the onset of an affective disorder results from the additive effects of several genetic and environmental factors. This concept is graphically illustrated in Figure 69.1. The horizontal axis represents the level of genetic predisposition, while the vertical axis represents the level of environmental risk. Thus, the more factors present (the number of implicated genes or environmental events that contribute to an affective illness), the more likely an individual would develop an affective illness and the more severe that illness would be. One could also assume that early-onset illnesses probably result from greater genetic loading, while late-onset illnesses may have less genetic loading but increased occurrence of environmental events with age.

The second explanation, increased phenocopies, illustrates the limited specificity of the affective disorders diagnosis. Both bipolar disorder and major depressive disorder are syndrome diagnoses, that is, the diagnosis is based on the presence of a collection of observed symptoms (phenotype) rather than an etiological cause. In this theory, late-onset affective illness is a collection of phenocopies. Phenocopies are illnesses with a

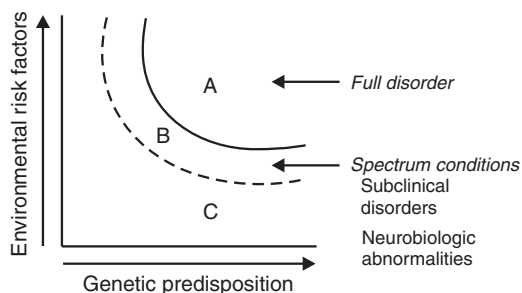


Figure 69.1 How genes and environment lead to illness. Adapted from Faraone *et al.*³¹

characteristic phenotype but different underlying causes. Hypothyroidism is an example of a phenocopy. Individuals with hypothyroidism may meet major depression criteria, but they require quite different treatment.

It should be noted that the two explanations postulated for the observed differences in age-related affective disorders are not mutually exclusive. We have already noted that affective disorders are heterogeneous. Figure 69.2 demonstrates two dimensions of heterogeneity—clinical and causal. Clinical heterogeneity occurs when more than one clinical condition can occur by the same cause. Causal heterogeneity occurs when two or more causes can independently produce the same clinical syndrome³¹. In Figure 69.2 the small circle at the top represents all patients who have the full disorder (in this case, bipolar disorder or major depressive disorder). The larger circle demonstrates three outcomes attributed to disease genes: “full disorder”, “spectrum conditions” and “symptom-free”. Some individuals with the disease genes will develop the full disorder. Others will not meet criteria for the full disorder, but will show abnormalities and trends toward the disorder. These are called “spectrum conditions”. Finally, some individuals will remain symptom-free despite carrying the same disease genes.

The small circle at the top of Figure 69.2 also shows that not all patients who have the full disorder have the disease genes. This

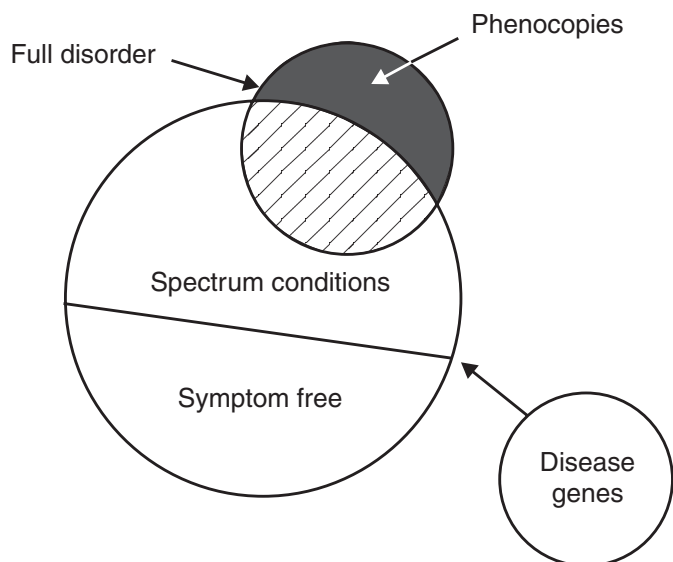


Figure 69.2 Causal and clinical heterogeneity. Adapted from Tsuang and Faraone¹¹

subgroup represents those patients with phenocopies of the full disorder. They have an illness that looks like a genetic disorder but it is not caused by genes, at least not the same disease genes.

It has long been known that depressive syndromes are very common in neurological disorders such as Parkinson’s disease, Alzheimer’s disease and stroke. More recently, researchers^{32,33} have proposed a subtype of depression called “vascular depression”, based on the presence of white and gray matter changes observed on magnetic resonance imaging (MRI) scans. These changes are frequently seen as patients age, and both the pattern and intensity of lesions appear to have a direct relationship to major depressive episodes (and possibly bipolar disorder). This subtype is differentiated from other depressive types by demonstrating a constellation of depressive symptoms, increased incidence of apathy, psychomotor retardation, cognitive impairment, functional disability and a decreased incidence of familial affective disorders. If this is shown to be true, then the age-related variation in late-life affective disorders may be explained by increasing phenocopies in older adults who have late-onset affective disorders. Patients with early-onset affective disorders (which would include elderly patients with previous diagnosis of affective disorders) may be more likely to have a genetic cause.

As research like this sharpens our understanding of disease, the “heterogeneous” group of affective disorders will be differentiated into various subtypes, each with a more specifically defined course, prognosis and treatment. Each new subtype will also suggest new ways that genes and the environment may contribute to the expression of the illness. For example, current research is examining how other “disease genes” may cause various phenocopies. One of the major complications of Alzheimer’s disease is the development of a major depressive disorder. Research has suggested that the presence of one of the ApoE alleles, which has been implicated in Alzheimer’s disease and is genetically determined, may predispose patients to the development of major depression^{34,35}. Other researchers have not found this finding to be consistent³⁶⁻³⁸. However, the implication is clear. Environmental and genetic patterns in late-onset affective disorders are emerging that are different from patterns seen in early-onset disease.

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Environmental Factors, Life Events and Coping Abilities

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Understanding environmental factors, life events and coping abilities in the lives of older people is best accomplished by applying a life-span developmental perspective. Baltes *et al.*¹ emphasize that people evolve throughout their lifetime and become more heterogeneous with age. As people grow older, the experiences of a lifetime combine uniquely to shape them. This is especially important to the question of how individuals cope with the challenges of old age. The lifespan framework allows us to incorporate information about an individual's experiences, the successes or failures an individual has had in coping with these experiences, and the people who have or have not been helpful in aiding the individual to achieve those successes. Social support is not always or solely considered a coping ability or strategy. However, it has been argued that successful coping is the ability to activate the kinds of support needed to meet the needs of specific situations and to help the individual achieve a positive affective disposition.

The Convoy Model of Social Relations^{2,3} suggests that individual and situational characteristics combine to explain and predict the amount, type and adequacy of support an individual is capable of garnering to cope with life events and circumstances, thus enabling the individual to adapt successfully to life's challenges. The individual moves through time and circumstance surrounded by a "convoy" of people, who either help or inhibit the individual's ability to successfully cope with the problems he/she faces, consequently affecting both his/her health and well-being. The Convoy Model incorporates both personal and situational factors to predict coping skills and success. Available evidence indicates that most convoys consist of both family and friendship relationships within a multigenerational context.

These contexts are characterized by exchanges that are perceived as either reciprocal or non-reciprocal. Since reciprocity is the norm of social relationships⁴, it is useful to consider how people assess the reciprocity of their relationships. As one gets older, it is often the case that one requires more support, both emotionally and instrumentally, to cope with the challenges of life. There is some danger that older people will come to perceive their relationships as non-reciprocal, and feel that they are over-benefited, i.e. receive more support than they provide, by these exchanges. There would then be the danger that this will lead to a sense of indebtedness. However, many people are quite resourceful about maintaining relationship equity by using what has been termed a "support bank"³ accounting system. Older people have been known to consider support provided by them to others earlier in life as contributions to a "support bank" from which they can make withdrawals, as needed, later in life. This

interpretation of support exchanges can be especially useful for understanding how some older people cope with crises by accepting considerable support from others but without the disadvantage of feeling overly indebted by the receipt of more aid than they are currently able to reciprocate. Also interesting is the phenomenon whereby the older person recalibrates support received of one type to equal support provided of another type. Thus, a drive to the doctor's office can be completely reciprocated by a freshly baked loaf of bread. Some might argue that the ability to cognitively construct an equitable exchange is not necessarily a reality assessment, but rather an exercise in adaptive illusions⁵.

Several recent theoretical developments should be noted. Baltes and Baltes⁶ have proposed the Selective Optimization with Compensation Theory. This theory suggests that an important strategy of successful aging is to carefully select those behaviors, goals or activities that are personally most important, and to develop multiple strategies to compensate for the limitations increasingly evident with age. This theory suggests that some goals will be ignored, dropped or otherwise disregarded, while others will be achieved but only through compromise and compensation. A related theory has been proposed, the Socio-emotional Selectivity Theory⁷. This theory focuses specifically on social relations and suggests that, with age, people reduce the number of social relationships in which they invest time, energy and emotional commitment, to only those which are truly close and important to them. The argument suggested is that with fewer years left to live, older people are less interested in relationships that are troublesome or otherwise unsatisfying to them and more interested in maximizing their time and commitment to those relationships that are truly important to them. It could also be that individuals are selected who allow the elder to maintain reciprocity or those with whom the elders have had a particularly satisfying exchange relationship in the past. Empirical data are available to support both perspectives. Especially interesting, in light of the Socio-emotional Selectivity Theory, is the finding that older people report significantly fewer negative social relations and are more satisfied than younger people with the relationships they do have.

We know that people vary in their ability to cope with the problems they face and in their ability to develop supportive relationships. We can best understand these differences as indications of how older people differentially experience their environments and life events and how they cope with these experiences. Although it is clearly true that some people experience more crises than others, it is not always the case that those who experience a great many crises are the ones

overwhelmed by them. The same environmental conditions or life events that might devastate one person may have less of a negative impact on another.

The Support/Efficacy Model⁸ proposes a lifespan framework to explain these differences. This model suggests that supportive interactions over time are essential to developing in the individual a sense of efficacy that instills in that person the belief that he/she can successfully meet the challenges of life. With time, exchanges and interactions accumulate which either lead the individual to feel competent and capable of coping with the problems he/she confronts (i.e. efficacious) or, under less optimal circumstances, leaves the individual feeling overwhelmed by, and personally incapable of, coping with the problems and circumstances of life.

It is also important to consider the influence of non-psychiatric factors on the experience of events in old age and how one copes with stress. Jackson *et al.*⁹ have noted that racial, cultural and other sociodemographic factors fundamentally affect how a situation is experienced. If an individual is one of many suffering the same negative experience(s), the etiology of that experience may not be devastatingly personalized. On the other hand, if everyone else in one's reference group is significantly more successful, the relative comparison can be devastating, even if "objectively" the situation is quite positive. Similarly, there now exists empirical evidence indicating that different national groups appear to vary considerably in the degree to which they respond negatively to environmental factors and life events. Fuhrer *et al.*¹⁰ have shown that normal French elderly score markedly higher on measures of depressive symptomatology than their American counterparts, despite the fact that their general environmental circumstances and their experienced life events are actually quite similar. Antonucci *et al.*¹¹, comparing Black and White Americans with French elderly on a variety of factors, have shown that the general mental health of American and French elderly are similar in some respects and different in others. All three groups are negatively affected by functional limitations. On the other hand, Black American and French elderly men are less satisfied with life than women in these groups, although there is no sex difference among White Americans. A recent study of the elderly in the Netherlands and Italy¹² is also of interest. It suggests that objective characteristics, such as living arrangements, which some might consider indicators of coping, need to be understood in terms of the individual's psychological state. Dutch elders tend to live alone, while Italians tend to live with their families. However, Italian elders reported less social integration and more loneliness than Dutch elders. Despite the appearance of family integration, these data suggest that an individual's ability to cope with environmental factors is not always directly predictable from the objective characteristics of the situation. As with the French and Americans reporting of depressive symptomatology, this could simply, but importantly, be a difference in expressive style.

It has been suggested that an individual's ability to cope with specific environmental conditions and life events is best understood through a consideration of the resources and experiences available to that person. One simplistic way to consider this is that successfully coping with stresses and strains in early life is the best predictor of the individual's ability to cope with stressful situations in later life. This is likely to be true, even if the exact nature of the stressful event varies. There is also reason to believe that some individuals develop coping styles over their lifetime that

can be seen to be generally successful and adaptive, while others develop coping styles that are generally unsuccessful or maladaptive. At best, these coping strategies match the environmental conditions and life events that the individual experiences. For example, Jackson¹³ and Jackson *et al.*¹⁴ have argued that the harsh environmental conditions faced by many racial and ethnic minorities lead to the development of novel and effective coping strategies. Over the lifecourse, however, many of these strategies actually may be deleterious to health status, e.g. alcohol use. Although the specific nature of these experiences will change with age, an individual's ability to cope with these environmental events is likely to show fairly stable lifespan continuity. At the same time, as research clearly demonstrates, individual coping and adaptation competencies can be improved through informal and professional intervention at all points in the individual lifecourse.

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The Aetiology of Late-life Depression

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It is vain to speak of cures or think of remedies, until such time as we have considered of the causes (Galen)

Empirics may ease, and sometimes help, but not thoroughly root out . . . as the saying is, if the cause be removed, the effect is likewise vanquished (Burton, *The Anatomy of Melancholy*)

Late-life depression, when defined according to the broad criterion of clinical significance, is a common disorder affecting 10–15% of the 65+ population^{1,2}. Prevalence rates for major depression are substantially lower, but this category excludes common forms of late-life depression, particularly those associated with bereavement and physical co-morbidity. Longitudinal population-based studies suggest that incidence and maintenance rates are both high, balanced by a high mortality ratio for those affected.

The aetiology of late-life depression remains unclear in some respects. While many population-based studies have been carried out in Europe and the USA, most of these have been cross-sectional in design, and have limited themselves to univariate analyses. The results of these investigations therefore appear as lists of cross-sectional associations, possibly confounded by other factors. While causal models can, and have been, inferred from such data, the process is fraught with difficulty and can lead to errors. More recently, good quality prospective studies have been carried out that have the potential to clarify the aetiology of these common disorders and inform primary and secondary prevention.

AGE

The data on the prevalence of major depression from the US Epidemiological Catchment Area (ECA) survey suggested a lower rate for those aged 65 (1.0%) than for those aged 45–64 (2.3%) and those aged 18–44 (3.4%)³. This relatively low prevalence rate among the older population was confirmed in a Canadian study using similar methodology⁴. These findings have been particularly controversial, as they could be taken to imply that the management of depression may require less resources per capita for the old than for the young⁵. They conflict with the general impression that the frequency of depressive symptoms and broader depressive syndromes either increases^{6–8} or remains stable⁹ with increasing age. The lay administered Diagnostic Interview Schedule (DIS) used to derive DSM-III diagnoses in the ECA excludes symptoms attributable to bereavement, physical illness or cognitive impairment. The ECA findings have been criticized on the grounds that the complex standardized symptoms, and the judgemental process required for responding to probes used in the DIS, may exceed the cognitive capacity of many older adults, leading to a systematic

response bias. This may have been a particular problem where subjects were required to attribute symptoms to physical or non-physical causes. Older subjects report as many lifetime depressive symptoms as younger subjects, but are more likely to attribute them to physical causes, meaning that they are then excluded as a basis of diagnosing depression¹⁰. However a re-analysis of ECA data reattributing physical symptoms to psychiatric symptoms did not lead to a disproportionate rise in major depression among the older age groups¹¹.

A further curiosity has been the consistent finding that the lifetime prevalence of major depression seems to be lower for elderly subjects (1.4% for those aged 65 in the ECA survey) than for younger subjects (7.5% for those aged 30–44, from the same survey). It has been suggested that this may represent a cohort effect, with successive birth cohorts carrying an increasing propensity for major depression. More plausibly, this finding may have arisen from a selective tendency for older subjects not to recall earlier undiagnosed episodes¹² and from the selective mortality of those most vulnerable to repeated severe episodes of depression^{8,13}. A broad review of this area reported similar findings for most psychiatric diagnoses, including schizophrenia, and concluded that cohort trends cannot be safely extrapolated from cross-sectional data¹⁴.

GENDER AND MARITAL STATUS

One of the clearest and most reproducible findings in psychiatric epidemiology is the apparent excess of depression among women. It has been suggested that the extent of this excess varies across the life course, increasing from menarche into mid-life, and then declining gradually into late life¹⁵. The EURODEP consortium^{16,17} reported a clear-cut excess of depression symptoms in older women in population-based studies from 13 out of 14 European centres. Interestingly, this association was consistently modified by marital status, with marriage being protective for men but a risk factor among women. Marriage is associated with relatively low mortality and good health, although this protective effect seems to be stronger for men than for women^{18,19}. In younger people, marriage also protects against depression among men but not among women²⁰. In Gove's²⁰ study, the excess of depression in women relative to men was greatest in married people. This striking finding has been variously attributed to the mundanity of housework and the unfavourable position of women who work outside the home²⁰ to the differences in the number and range of role identities by gender and marital status²¹ and to the burden of childcaring²². Brown *et al.* drew attention to the lack of satisfaction with the married state expressed by

working class Camberwell women. In older people these observations are consistent with the observation from several studies that married older men cite their wife as their main confidante, whereas women more often cite a friend outside the home²⁴. Also, in Finland, a prospective study showed that for men the risk of onset of depression over 5 years is increased for those having poor emotional relations with their wives, while for women the risk is greatest among those not living alone at the beginning of the follow-up period²⁵. These findings led Kivels to suggest that marital counselling should be made available for older people. However, there may be external factors which, to the extent to which they affect wives and husbands and single men and single women differently, may have explained some of the observed gender/marital status interaction. One such factor may be the social integration and activity of single men and women. Never-married men reported fewer supportive friends and neighbours, less attendance at clubs or church, and more loneliness than never-married women in Gospel Oak^{26,27}. Another area worthy of investigation is the relative health of male and female marital partners. A national US survey showed that 64% of all spousal carers were wives, suggesting that in older age the burden of care in marriages may generally derive from the husband and devolve on the wife²⁸.

DISABLEMENT

Of the world population, 7–10% are significantly affected by disablement, defined as the long-term consequences of chronic disease²⁹. These estimates have varied little between world regions, and are similar for developed and developing countries²⁹. There is, however, a strong positive relationship between disablement and age. The OPCS Surveys of Disability in Great Britain reported a near doubling of disability prevalence rates with each 10 year increase in age, from 7% for those aged 40–49, to 67% for those aged 80+³⁰. Among older subjects, most detectable physical illness is chronic rather than acute³¹.

Many studies have commented on the strength of the cross-sectional relationship between physical health variables and depression in older age. Gurland reviewed 70 years of research endeavour in this field, carried out in primary care, hospital and community settings, and concluded that there was strong accumulated evidence of a tendency for both physical illness and disability to co-exist with major and minor depression at a frequency greater than that expected by chance³².

There have been suggestions from clinical populations of specific associations between late-life depression and diseases such as stroke³³ and Parkinson's disease³⁴. Results from community surveys are less clear. However, stroke, respiratory disease and arthritis were all found to be associated by most studies which have assessed individual diagnoses^{35–40}, while hypertension and diabetes were less salient. This pattern of association with individual diseases would suggest that disablement, the limiting long-term consequences of disease, may be more relevant than any particular pathology. Indeed, the strongest reported associations have generally been between depression and summary measures of disablement. A systematic review of the literature, using MEDLINE for the period 1984–1996 and secondary references for earlier publications, revealed 10 cross-sectional studies which had used population samples⁴¹. They were consistent in reporting strong positive associations between disablement, measured in various ways, and depression^{36–38,42–46}. However, the strong associations might not reflect a causal relationship. Bias (somatic contamination of the measurement of depression), confounding and, in particular, reverse causality (depression leading to disablement, rather than vice versa) were plausible alternative explanations. Later prospective population-based research has

clarified the association. At least five longitudinal studies have now shown a very strong association between disablement at baseline and the subsequent onset of depression^{47–51}. In the Gospel Oak survey^{26,27}, after adjusting for confounders, the most restricted quarter of the Gospel Oak population (London Handicap Scale) were 20 times more likely to be depressed at baseline than the least restricted quarter. Those among them who were not depressed at baseline were five times more likely than the least restricted quarter to have experienced an onset of depression at 1 year⁵⁰. The population attributable fraction (the proportion of new cases that might notationally be prevented if the risk factor were removed) was 0.69. Most studies agree that it is the level of disablement associated with a health condition, rather than the nature of the pathology, that determines the risk for depression^{50,52,53}. Three population-based studies have suggested an interaction between disablement and social support, with the strongest effect of disablement in those with the least social support^{1,50,51}. Beekman *et al.*¹ reported that the association between disablement and depression was only apparent for minor rather than major depression. However, this finding was based upon cross-sectional research, and requires replication. Interestingly, in community studies, disablement does not seem to influence directly the process of recovery from late-life depression⁵⁰.

Ormel *et al.*⁵⁴ argue in a separate publication that synchronicity of changes in depression and disability observed longitudinally in a primary care-based study support the hypothesis that there is an important pathway leading from depression to disability. In reality, as most authors have acknowledged, the situation may be more complex; the causal direction may vary between individuals, and components of each direction may co-exist within the same individual. A case can also be made for reciprocal causation, with a physical impairment leading to handicap, provoking depression which may in turn exacerbate the degree of handicap associated with the original impairment.

LIFE EVENTS

The literature on life events in older people was recently reviewed⁵⁵. In the main, two methods have been used. The Bedford College Life Events and Difficulties Schedule (LEDS)⁵⁶ elicits events in a lengthy semi-structured recorded interview. These are then rated independently for contextual threat by a trained panel. Studies using this detailed method have shown that depressed older subjects have experienced more recent life events than non-depressed subjects⁵⁷. However, older samples differ from younger ones in that chronic difficulties are more prevalent than life events, and events typically carry relatively low levels of threat^{58,59}. Also, in contrast with Brown's work on younger adults, health difficulties are an important source of adversity^{57,58}. The second and more common approach has been to use a pre-determined checklist of events. In a community survey in the USA, the onset of illness affecting a subject or relative, the onset of money problems, and becoming a victim of crime were among the most common and most undesirable life events affecting subjects aged 55+⁵⁹. Deaths of spouses, children and siblings were rated as highly undesirable, but were individually relatively infrequent. However, 14% of females and 12% of males experienced a family bereavement of some kind over 1 year. Most checklists focus on these event categories, with a particular bias towards bereavement and personal illness events. Evidence for a relationship between depression in older age and life events measured using checklists is generally weaker than in LEDS-based studies. Linn⁶⁴ found a small significant difference in the mean number of events experienced by depressed and non-depressed community subjects. Other studies have not replicated this finding^{60–63}. In this study,

subjects' history of exposure to serious life events was ascertained retrospectively, at index assessment, over a period of up to 2 years prior to the time of interview. Perhaps for this reason, associations with prevalent depression at index assessment were more impressive than the prospective associations with the onset of depression 1 year later. It may be that the categories used by some life event checklists carry too little "contextual threat" to be relevant to the subsequent onset of depression. In the longitudinal Gospel Oak survey^{20,27}, at index assessment depressed subjects reported more life events, measured using a checklist of events likely to carry high contextual threat, than did non-depressed subjects. However, at follow-up only a weak non-significant trend towards an increased risk for the onset of depression was observed. Having something important lost or stolen was the only individual event associated with the onset of depression over 1 year.

SOCIAL SUPPORT AND THE BUFFER HYPOTHESIS

Brown demonstrated in a younger sample that negative self-esteem and lack of social support were vulnerability factors, increasing the risk of depression in the presence of a life event²³. Murphy⁵⁷ reported a similar vulnerability for older subjects lacking a confiding relationship. Not all studies have replicated this finding. Murrell⁶¹ found a direct protective relationship between self-esteem and depression, but noted that neither self-esteem nor health modified the association between life events and later depression scores. Social support has been infrequently investigated in older community samples, and the findings have not all been consistent. Murphy's finding of an association between a lack of confidants and depression was confirmed in New York⁴⁴. Bowling reported a positive association between the number of confidants and well-being in a London sample, which was, however, not apparent in a sample drawn from a smaller town in Essex⁶⁵. In both samples, health status accounted for more of the variance in well-being than did subjects' social support or social network characteristics. Bowling's finding of no association between social participation and well-being was contradicted by Palinkas' report, in a Californian study, of inverse associations between participation in church and other community organizations and depression^{35,36}. Palinkas also reported an inverse relationship between social network size and depression. Both Palinkas^{35,36} and Woo⁶⁶ in a study from Hong Kong⁶⁶, report inverse associations between the frequency of contacts with friends and depression. The Bowling, Woo and Palinkas studies all adjusted to some extent for subjects' physical health status, and Palinkas also controlled for age and gender. Each of these variables is a potential confounder of an association between social support and late-life depression. There are large differences between the social support networks of older men and women, women typically having more supportive and extensive networks of friends than men^{35,36,67}. Palinkas hypothesized a deterioration of social network with increasing age consequent upon bereavement; he found an inverse association between age and network size in women, but not in men. Bowling⁶⁵ comments that active social engagement, such as visiting friends, is often confounded by functional ability, but unfortunately does not report her data on associations between social support or social network and health status. Palinkas found no association between physical health and social participation, or the number or frequency of social contacts^{35,36}, but other studies have observed disablement to be associated with smaller social networks^{68,69}.

One of the more consistent findings from the literature appears to be the salience to late-life depression of contact with friends, in particular intimate, confiding relationships. While older people typically receive instrumental support from spouses and relatives,

they value friends for the companionship and emotional support that they can provide⁷⁰. A review of recent research in the area concludes that interaction with friends, rather than frequency of contact with relations, is positively associated with emotional well-being²⁴. This view was borne out in the longitudinal Gospel Oak study. No contact with friends was the only social support variable prospectively associated with the onset of depression⁵⁰. In this study, lack of social support and social participation were more evidently associated with the maintenance rather than with the onset of depression.

SOCIAL CLASS, INCOME AND EDUCATION

Brown and Harris reported that working-class women with children at home were four times more likely to have experienced a depression than middle-class women²³. Although they were more likely to experience life events, the class difference in depression was mediated more by the higher prevalence of four vulnerability factors among the working-class women. Murphy has reported a less dramatic social class effect in an older sample⁵⁷. In her older subjects, the excess of depression among the lower social classes was largely explained by their poorer health and greater social difficulties. Social class, as measured by best pre-retirement occupation, may retain some validity after retirement, to the extent to which it relates to retirement income, housing quality, social network, and access to and use of formal services. There have been reports from cross-sectional community surveys from a variety of cultures of associations between late-life depression and relative disadvantage in income^{37,38,44,66,71,72}, housing status^{37,38,66} and education^{37,38,44,46,66,72}. These are, of course, highly correlated variables, and it will always be difficult to determine the effect of one independent of the others. The possibility of reverse causality also needs to be considered because of the well-recognized phenomenon of social drift; people whose adult life has been scarred by depression may experience occupational and economic disadvantage.

FAMILY HISTORY, GENETIC LIABILITY AND PAST PSYCHIATRIC HISTORY

The earliest evidence of a familial tendency in depression came from family studies in which the life-time prevalence of depressive disorder was compared between relatives of probands with depression and a control sample. Many such studies have been completed, with consistent results. Higher than expected rates of major depression and minor depression are found among relatives of probands with major depression, and higher rates of bipolar disorder and major depression among relatives of probands with bipolar disorder⁷³. Such studies could only identify familial aggregation, and could not distinguish between familial similarity arising from shared environment or shared genes.

Early twin studies suggested a substantial contribution from both genes and shared environment⁷⁴. Incorporation of environmental measures in genetically sensitive designs has advanced our understanding of the interplay between genetic and environmental risk factors for major depression. Kendler *et al.*⁷⁵ have completed extensive quantitative genetic analyses in a large cohort of same-sex (female) twins from the population-based Virginia Twin Registry. The heritability for a life-time diagnosis of major depression was 33–45%, depending on the diagnostic criteria that were applied. This estimate varied little between more and less restrictive criteria⁷⁵. The heritability for 1 year prevalence of major depression also lay between 41% and 46%⁷⁶. There was some suggestion that improving the reliability of the diagnosis of major depression might increase the estimates for the heritability

of the disorder, with up to half of the environmental contribution to the variance having reflected measurement error, leading to a dilution of the previous reported genetic contributions⁷⁷. About 60% of the genetic effect contributed was indirect, mediated by past history of depression, neuroticism, life events and childhood adversity. Two coping strategies, turning to others and problem solving, which had been negatively correlated with levels of depression, were also found to have substantial heritability⁷⁸. The relationship between genetic liability, stressful life events and depression was explored in detail, and evidence provided to support a gene-environment interaction in which genetic factors determined sensitivity to serious life events⁷⁹.

Data on older subjects is relatively lacking. Two studies from The Netherlands examined cross-sectional relationships between depression and the exposures of a family history of mental illness, and a personal past history of depression. In the Amstel study, based on a large population sample of 2540 subjects aged 65+, there was a positive association between a family history of mental problems and a diagnosis of GMS-AGECAT depression⁸⁰. This association was modified by both cognitive status and personal past psychiatric history. The strongest association was seen in the group with a past psychiatric history and no cognitive impairment. However, only 22% of depressed subjects reported a family history of mental health problems, suggesting a modest population-attributable fraction. Van Ojen¹³ also reported that, while depression was twice as common in those with a psychiatric history with an onset before the age of 65, the prevalence of this exposure decreased linearly with increasing age¹³. Only 22% of currently depressed subjects reported a previous episode of psychiatric illness with an onset before age 65. While failure of recall and "telescoping" of ages of onset might have accounted for these findings, the reduction in the prevalence of past history with increasing age was consistent with predictions based upon the known excess mortality among those so affected. The LASA study, based on a national Dutch sample of over 3000 subjects aged 65+, used two definitions of depressive disorder, major depression diagnosed using DIS, and minor depression defined as those scoring 16 or more on the Center for Epidemiological Studies Depression Scale. A family history of depression was associated with neither outcome. A past history of psychiatric disorder was strongly associated with major depression, with an odds ratio (OR) of 90.8 (39.1-211.1) and a population-attributable fraction (PAF) of 74%, but much more weakly associated with minor depression; OR 4.1 (2.3-7.3) PAF 7%.

COGNITIVE DECLINE

An inverse association between depression scores and cognitive test scores has been reported in four cross-sectional surveys^{46,71,81,82} with one negative report⁴². These cross-sectional studies did not permit further analysis of direction of causality. However, at least one study has suggested that low mood may be a risk factor for subsequent decline in cognitive ability⁸³. A biological basis for this association has been suggested; the failure, seen in chronically depressed subjects, of fast-feedback mechanisms to shut off the cortisol stress response may lead to high concentrations of neurotoxic metabolites, causing specific damage to hippocampal regions essential to new learning and memory recall tasks⁸⁴. Alternatively, microvascular lesions leading to cognitive decline may also cause late-life depression through critical subcortical damage to frontotemporal cortical projections necessary for the maintenance of mood and motivation. MRI evidence suggests that first onsets of late-life depression may be associated with characteristic deep white matter lesions⁸⁵. The finding from the Amstel study, that the associations between life events and depression and living alone and depression were not

apparent in those with cognitive impairment, would be consistent with a hypothesis of an alternative pathway to late-life depression mediated through cerebral deterioration⁸⁶. There may be a more prosaic explanation for a prospective association between cognitive decline and depression. Cognitive impairment, like other health impairments, may lead to depression to the extent to which it causes disability and handicap.

CONCLUSION

The factors most consistently implicated in these studies are similar to those observed in younger adults: constitutional or genetic vulnerability, e.g. a family history or past history of psychiatric disorder; current vulnerability, e.g. lack of social support, recent stress, as in life events, and current adversity, as in low socioeconomic class, poor housing or low income. However, some factors may be particularly salient for late-life depression, either because, as in the case of disablement or bereavement, they are a much more common exposure among the older population, or because they may impact differently upon those who are exposed, depending on their age. There is already evidence to suggest that disablement associated with declining health in older age may be a prime determinant of the prevalence, incidence and maintenance of late-life depression.

In younger adult populations, depression is a substantially heritable disorder in which both direct genetic effects and genetic effects mediated through childhood adversity, neurotic personality, coping styles, propensity for life events and sensitivity to life events can be discerned. Evidence from genetically sensitive designs in older populations is lacking, but circumstantial evidence suggests that the selective early death of those with an early first onset and multiple episodes of depression, who perhaps have a higher genetic loading for the disorder, may mean that the heritability of depression in late life is lower than in younger populations. The finding that the large majority of older depressed subjects are experiencing a first onset of the disorder raises the possibility of aetiological dissimilarity between early adult and late-life depression. There is a clear case for focusing in our investigations on those aspects of physical health status, cognition and social milieu that change most acutely in late-life and best distinguish the life experience of older and younger adults.

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Risk Factors and the Incidence of Post-stroke Depression

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University of Western Australia, Australia

Post-stroke depression (PSD) is the most common psychiatric syndrome following stroke, but there is no specific PSD syndrome. It is commonly unrecognized and untreated in clinical practice. Rao¹ has reviewed the literature on the relationship between depression and cerebrovascular disease, including strokes.

PREVALENCE

Reported prevalence of PSD has varied between 30–60%, 8–30% major and 20–40% minor depression. The reasons for this wide variation include different assessment methods, diagnostic criteria, time elapsed after the stroke and patient settings, together with assessment difficulties in the presence of aphasic, cognitive and physical disabilities.

The lowest number of reported cases have come from community population-based studies. Several authors have concluded that it is unproved that depression is commoner in

an unselected group of patients after strokes than it is among the elderly with other physical conditions. Most cases of PSD develop during the first 2–3 months post-stroke.

RISK FACTORS

Described pre-stroke risk factors include personal or family history of depression or other psychiatric disorder, socially impaired personality, neuroticism, high alcoholic intake, social isolation and negative life events. Post-stroke factors include perception of the effects of the stroke as a significant loss, dysphasia, significant disability, impaired cognition, living in a nursing home, isolation and poor social supports. Most, but not all, studies have reported age, gender, educational level and marital status as unimportant.

Described biological risk factors include pre-stroke subcortical atrophy, especially of periventricular white matter, site and size of the lesions, and ischaemic rather than haemorrhagic stroke.

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an unselected group of patients after strokes than it is among the elderly with other physical conditions. Most cases of PSD develop during the first 2–3 months post-stroke.

RISK FACTORS

Described pre-stroke risk factors include personal or family history of depression or other psychiatric disorder, socially impaired personality, neuroticism, high alcoholic intake, social isolation and negative life events. Post-stroke factors include perception of the effects of the stroke as a significant loss, dysphasia, significant disability, impaired cognition, living in a nursing home, isolation and poor social supports. Most, but not all, studies have reported age, gender, educational level and marital status as unimportant.

Described biological risk factors include pre-stroke subcortical atrophy, especially of periventricular white matter, site and size of the lesions, and ischaemic rather than haemorrhagic stroke.

Disruption of adrenergic and serotonergic pathways by the lesion, giving depletion of biogenic amines, has been postulated.

Robinson and Starkstein² considered the location of the lesion in the brain as the single most important aetiological factor in PSD, with the highest frequency of PSD associated with anterior lesions in the left hemisphere. Their method of neuroradiological lesion location has been criticized³. Lesions of the globus pallidus, putamen and caudate nucleus in association with PSD have been described.

A recent literature review by Singh *et al.*⁴ concluded that any definitive statements about stroke lesion location and risk of depression have not yet been substantiated. Anatomical correlates of PSD change over time and may explain interstudy differences in the association of lesion location with PSD⁵. Furthermore, mood changes may be mediated through the distant effects of lesions, as shown by PET studies, or the lesions may play a facilitatory role for another primary mechanism leading to depression. Many of

these risk factors may be interrelated, e.g. size of lesions, disability, impaired cognition and living in a nursing home.

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Epidemiology of Depression: Prevalence and Incidence

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Many symptom checklists have been used to estimate the burden of depressive symptoms in community populations. The results of these studies are remarkably consistent, with the range of significant depressive symptoms estimated to be 10–25% (see Table 71.1). Estimates of the prevalence of major depression and dysthymic disorder are presented in Table 71.2 from four different countries. The prevalence of major depression is much lower in community samples than is the prevalence of depressive symptoms, 1–5% in most community surveys. Lower prevalences of major depression have been found in more rural samples with an estimate of less than 1% in a North Carolina rural sample, whereas urban studies have typically estimated higher prevalence. The prevalence of dysthymic disorder is generally higher than that for major depression, approximately 2–8%, depending on the instrument used in the survey. Prevalence estimates are difficult to compare cross-nationally, as different instruments are used. Comparison is even more difficult when symptom burden is confused with diagnosed cases. For example, if the prevalences of dysthymic disorder and major depression are combined and the frequent comorbidity of dysthymic disorder and major depression taken into account, the majority of subjects suffering significant depressive symptoms in community populations still do not qualify for diagnosis. One of the major tasks facing psychiatric epidemiologists studying depression cross-nationally in the elderly is to explain the residual depressive symptoms in community samples not easily captured by the usual diagnostic categories.

In a recent review of 34 studies of the prevalence of late-life depression by Beekman *et al.*¹¹ (part of the EURODEP Study),

the prevalence varied (0.4–35%). Arranged according to diagnostic category, major depression was relatively rare (weighted average prevalence of 1.8%), minor depression more common (weighted average prevalence of 9.8%) while all depressive syndromes deemed clinically relevant yield an average prevalence of 13.5%. Depression was more common among women and among older people living under adverse socioeconomic circumstances. In a cross-national comparison of nine European centers using the GMS-AGECAT package to abstract diagnoses, differences in prevalence by country were most evident, with a range from 8.8% in Iceland to 23.6% in Munich. The overall prevalence estimated by meta-analysis was 12.3%, 14.1% for women and 8.6% for men. Among the elderly, there was little variation by age across all sites.

The prevalence of depression in acute medical facilities is presented in Table 71.3. Estimates range from 5% to 10% for major depression but, of more importance, an additional 15–25% experience clinically significant symptoms not captured by the diagnosis of major depression. The same is true for estimates of prevalence in long-term care facilities (see Table 71.4). Estimates range between 5% and 15% for major depression, yet an additional 30% of the sample subjects (in one study) suffered clinically significant depressive symptoms that did not meet the criteria for major depression. Estimates of depressive symptoms and depressive diagnoses (major depression and dysthymic disorder) among the elderly should not be based solely on community samples. Older adults in hospital or residing in long-term care facilities are much more likely to

Table 71.1 Prevalence of depressive symptoms in community samples of older adults

Author	Sample	(n)	Screening method	Findings
Blazer and Williams, 1980 ¹	Community sample in North Carolina (65+)	997	OARS Depression Scale	14.7% significant depressive symptoms. No age, gender or racial difference in prevalence
Kivela <i>et al.</i> , 1986 ²	Community sample of elderly in Finland	1529	Zung SDS	29.7% of females and 22.4% of males with significant depressive symptoms
Ben-Arie <i>et al.</i> , 1987 ³	Community sample of the elderly "coloured" in Cape Town	139	PSE symptoms	13% with significant depressive symptoms
Kennedy <i>et al.</i> , 1989 ⁴	Community sample in Bronx, NY	2317	CES-D	16.9% with significant depressive symptoms, more prevalent in females and the oldest-old
Ihara, 1993 ⁵	Rural community sample in Japan	695	CES-D	5.3% with significant depressive symptoms
Livingstone <i>et al.</i> , 1990 ⁶	Community sample of elderly in inner London	813	CARE	15.9% with significant depressive symptoms

OARS, Older Americans Resources and Services; SDS, Self-rating Depression Scale; PSE, Present State Examination; CES-D, Center for Epidemiologic Studies Depression Scale; CARE, Comprehensive Assessment and Referral Evaluation

Table 71.2 Prevalence of dysthymia, minor depression and major depression in community samples of older adults

Author	Sample	(n)	Screening method	Findings
Blazer <i>et al.</i> , 1987 ⁷	Community sample in North Carolina, USA	1304	Diagnostic Interview Schedule (DIS)	0.8% with major depression, 2% with dysthymia, 4% with minor depression
Copeland <i>et al.</i> , 1987 ⁸	Community sample in Liverpool	1070	Geriatric Mental State Schedule	2.9% with major depression, 8.3% with minor depression
Beekman <i>et al.</i> , 1995 ⁹	Community sample in The Netherlands	3056	CES-D/DIS	2.0% with major depression, 12.9% with minor depression
Pahkala <i>et al.</i> , 1995 ¹⁰	Community sample in Finland	1086	DSM-III diagnosis	2.2% with major depression, 14.3% with minor depression

Table 71.3. Prevalence of depression in acute care medical facilities

Author	Sample	(n)	Diagnostic method	Findings
Koenig <i>et al.</i> , 1986 ¹²	VA inpatient sample of men 70+	171	Screening plus modified DIS	11.5% with major depression; 23% with significant depressive symptoms
O'Riordan <i>et al.</i> , 1989 ¹³	Acute medical geriatric assessment unit	111	Geriatric Depression Scale and Clinical Interview	4.5% with major depression; 3.6% with dysthymic disorder; 10.8% with significant depressive symptoms (most with dementia)

experience depressive symptoms and be diagnosed with major depression or dysthymic disorder than persons living in the community.

Incidence studies of depression in the elderly are extremely rare in the literature. Two studies provide some estimate of incidence, however. Rorsman *et al.*¹⁸ estimated incident depression from the Lundby cohort in Sweden among 2612 individuals evaluated in 1957 and later in 1972 (i.e. 15 year incidence) until the age of 70. The cumulative probability of suffering a first episode of depression was 27% for men and 45% for women, a very high incidence figure in this cohort (especially compared to lifetime prevalence figures, reported in other studies, of less than 15%). The annual age-standardized first incidence for depression, all degrees of impairment included, was 0.43 for men and 0.76 for women. Incidence appears to decrease in the studies as individuals aged, especially for men. Eaton *et al.* (1997)¹⁹ estimated the incidence of major depression over 10 years for the ECA cohort from Baltimore. They found an overall estimated annual incidence of 3.0 per 1000 per year, with a peak while subjects were in their 30s, a smaller peak when subjects were in their 50s and a definite lower incidence in the elderly. Prodromal symptoms were present many years before the full criteria for major

depression were met, further linking the minor and major depression, Foster *et al.* (1991)²⁰ estimated the incidence of depression in long-term care facilities. In a cohort of 104 new admissions followed for a year, they found an incidence of 14%. One-third of these new cases were diagnosed as major depression, two-thirds as minor depression.

HISTORICAL TRENDS OF DEPRESSIVE DISORDERS

Historical studies in epidemiology assist investigators to establish the frequency of disorders in populations at different points in time. To understand the prevalence of major depression in late life compared to earlier stages of the life cycle in modern Western societies (e.g. major depression appears to be less frequent in older adults), an historical approach is necessary. Depressive disorders, such as tuberculosis, acquired immune deficiency syndrome (AIDS) and smallpox, wax and wane in frequency through time. Unfortunately, historical studies in psychiatric epidemiology are rare. Therefore, temporal changes in mental illness are difficult to determine.

Table 71.4. Prevalence of depression in long-term care facilities

Author	Sample	(n)	Diagnostic method	Findings
Parmelee <i>et al.</i> , 1989 ¹⁴	Nursing home and congregate apartment residence in Philadelphia	708	DSM-III-R checklist	12.4% with major depression; 35% significant depressive symptoms
Bond <i>et al.</i> , 1989 ¹⁵	Three British NHS nursing homes	568	Crichton Royal Behavioral Scale and the Survey Psychiatric Assessment Scale	32–42% with severe affective disorder or psychoneurosis
Phillips and Henderson, 1991 ¹⁶	24 Australian nursing homes	323	DSM-III-R criteria	6.1% of residents suffered from a severe depressive episode, 6.7% from a moderate depressive episode and 6.7% from a mild depressive episode
Gerety <i>et al.</i> , 1994 ¹⁷	5 Nursing homes	135	Structured Clinical Interview for DSM-III-R diagnoses	26% diagnosed with a major depressive episode

Some investigators, however, have explored historical trends via the so-called "cohort effect". Klerman²¹ noted that, in contrast to the "age of anxiety" following the Second World War, modern Western society may be entering an "age of melancholy", precipitated by social factors, such as the threat of nuclear warfare, a perceived threat of environmental pollution and economic instability. Estimates of prevalence by age from the Epidemiologic Catchment Area Studies in the USA reveal a significant decrease in both current and lifetime prevalence of major depression by age cross-sectionally. In a separate study, Klerman *et al.*²², analyzing family history data from subjects in the Psychobiology of Depression Study, found a progressive increase in the rates of depression in successively younger birth cohorts throughout the twentieth century, with an earlier age of onset of depression with each successive birth cohort. However, these cross-sectional studies, which suggest an "age of melancholy", must be considered in the context of historical studies.

To interpret historical trends, one must consider the cohort effect. Four findings suggest that younger birth cohorts carry a higher prevalence of depression than older cohorts. First, longitudinal studies have typically demonstrated a consistent burden of depression across the life cycle for each birth cohort²³. That is, the prevalence of depression within a birth cohort does not change with increasing age. Second, younger birth cohorts appear to experience higher prevalences of depression than the older birth cohorts in the latter two decades of the twentieth century^{24,25}. Third, suicide rates do increase with age, which in turn drives the overall rate of suicide up with age²⁶. Fourth, the changes in rates of suicide and depression by age group over the past 50 years have been of such magnitude that the varying rates are best explained by psychosocial rather than biological or evolutionary factors.

Those factors that contributed to a relative protection of older birth cohorts from depression and suicide in the 1990s in Western societies, compared to younger cohorts, and that contributed to the relative increase in depression and suicide among younger cohorts in the 1990s, are unknown (but subject to much speculation). Interpreting these data, investigators and clinicians must take care to recognize that significant methodological problems in historical studies remain. Even a clear and decisive endpoint, such as suicide, can be misleading, for suicide rates are obtained from a review of death certificates, which are subject to considerable bias. Methods for estimating the prevalence of depression change through time, and therefore it is difficult to compare studies from many years in the past with current studies (due to an evolving instrumentation for measuring depression in community and clinical samples).

CONCLUSION

In summary, it appears that results from cross-sectional studies in Western societies estimate the prevalence of depression in older adults to be lower than the prevalence in young and middle-aged adults. These findings are most striking in the USA (particularly the Epidemiologic Catchment Area Studies) but also appear in other studies. Nevertheless, these findings must be interpreted with caution. Methodological problems in identifying depression across the life cycle may contribute to these difficulties. In addition, birth cohorts, with the well-known stability of function and disease that birth cohorts demonstrate, must be taken into account before cross-sectional data are translated into longitudinal interpretations.

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Epidemiological Catchment Area Studies of Mood Disorders

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The National Institute of Mental Health Multi-site Epidemiologic Catchment Area (ECA)¹ Program consists of a combined community and institutional survey of five communities in the USA: New Haven, Connecticut; Baltimore, Maryland; Durham, North Carolina; St Louis, Missouri and Los Angeles, California. Because of the large sample drawn for this study and because oversamples of the elderly were drawn at three sites, data are available that permit the estimate of the prevalence of affective disorders from a larger sample of community-dwelling elders than from any other extant study.

The goals of the Epidemiologic Catchment Area Program were to: (a) estimate the prevalence of specific psychiatric disorders using a similar methodology across multiple geographic sampling areas; (b) determine correlates of these specific psychiatric disorders; and (c) determine the relationship between psychiatric disorders and health services utilization.

The 6-month prevalence of the affective disorders from the five ECA sites overall range from 4% to 7%². All affective disorders, except for bereavement, were less prevalent in the elderly than at other stages of the life cycle. For example, the prevalence of major depression in men ranged between 1% and 4% among the 18–24 year-olds but was consistently less than 1% in the 65 age group. Among 18–24 year-old women, the prevalence of major depression was 7%, whereas in women in the 65+ age group it did not exceed 3% at any of the ECA sites. Recent incidence studies from the ECA sample suggest that the incidence for major depression peaks in the 30s with a smaller peak during the 50s. Incidence is much lower for the elderly.

Dysthymic disorder varied less by age in prevalence than major depression and was more prevalent than major depression in the elderly³. For example, the prevalence of dysthymic disorder was 0.5–3% in 18–24 year-old men and 0.5–2% in 65 year-old men. Among women, dysthymic disorder was 1–4% in 18–24 year-old women and between 1–4% in 65+ year-old women. Overall, current affective disorders of all types were less frequent in older persons (65+ years of age) than for any other age group. Manic disorders were extremely rare in the sample overall (0.5–1%). No cases of mania were identified in the 65+ age group across the five ECA sites.

The lifetime prevalence rates for the DSM-III specific psychiatric disorders evaluated in the ECA sample paralleled

rates of current prevalence but, as would be expected, were higher³. Lifetime prevalence for major depression was 4–10% in the 25–44 age group but was not higher than 2% in the 65+ age group at any ECA site. Only one lifetime occurrence of manic episode was identified in this very large sample.

The ECA studies have received considerable criticism from geriatric psychiatrists who perceive that the study design significantly underestimates the prevalence of affective disorders in older adults. The dramatic differences in prevalence (both current and lifetime) by age surely calls for some explanation. Studies are emerging to suggest that the Diagnostic Interview Schedule, the instrument used to determine the prevalence of psychiatric disorders in the ECA sample, may possibly be biased toward underestimating the prevalence of affective disorders in older persons. Specifically, the threshold for a symptom being identified by the instrument may be increased for older persons. Nevertheless, the threshold effect does not appear to explain the dramatic differences in prevalence and has led a number of investigators to suspect that a cohort phenomenon is operative⁵. That is, older persons not only have lower current prevalence of depression in the 1990s, they have always experienced a lower prevalence. According to a number of studies, this is true among more modern, Western societies.

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EURODEP—Prevalence of Depression in Europe

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The aims of the EURODEP Concerted Action were to use existing studies which had employed the GMS-AGECAT method of diagnosis to assess the prevalence of depression in nine European countries (10 centres)—Liverpool, Amsterdam (sample A), Berlin, Dublin, Iceland, London, Maastricht, Munich, Verona and Zaragoza. Later Tirana was added (not reported here), and five non-AGECAT centres joined after the study commenced—Bordeaux, Oulu, Antwerp, Amsterdam (sample B) and Göteborg.

All subjects were aged 65 or over. In Munich the subjects were aged 85 and above and in Iceland 85–87. The Amsterdam study (sample A) had an upper age limit of 84. All the studies took random samples in the community except Dublin, which sampled a general practice. Sample size varied from 202 in Verona to 5222 in Liverpool, giving a total sample of 13 803. Substantial differences in the prevalence of depression were found, with Iceland having the lowest level at 8.8%, followed by Liverpool, 10.0%; Zaragoza, 10.7%; Dublin, 11.9%; Amsterdam, 12.0%; Berlin, 16.5%; London, 17.3%; Verona, 18.3%; and Munich, 23.6%. When all five AGECAT depression levels, including both subcases of depression and cases, were added together, five high-scoring centres emerged, namely Amsterdam, Berlin, Munich, London and Verona (30.4–37.9%) and four low-scoring centres, Dublin, Iceland, Liverpool and Zaragoza (17.7–21.4%). There was no constant association between prevalence and age. A meta-analysis of the pooled data on the nine European centres yielded an overall prevalence of 12.3% (95% CI, 11.8–12.9); for women, 14.1% (95% CI, 13.5–14.8) and for men, 8.6% (95% CI, 7.9–9.3%)¹.

The proportions of depressive symptoms were found to vary between centres. In Amsterdam, for example, 40% of a general population of older people admitted to depressive mood, compared to only 26% in Zaragoza. Symptoms such as “future bleak”, “hopelessness”, “wish to be dead”, were generally rare, but the last reached higher levels in Berlin, Munich and Verona. Sleep disturbance was admitted by only 15% of the population in

Dublin, but 54% and 60% in Munich and Berlin. Large differences for some depressive symptoms were found within the very old populations, with lower levels in Iceland and higher levels in Munich. Overall, the levels of depressive symptoms among over 60% of the older general population of Europe were low, so that pejorative stereotypes of old age in Europe were not upheld².

In order to include non-AGECAT centres, attempts were made to harmonize the depression measures that had been used with items from the Geriatric Mental State examination. A scale was constructed, the Euro-D scale³. The scale appeared to work well and was applied to data from 21 724 subjects. Euro-D scores tended to increase with increasing age, unlike the levels of prevalence of depression. Women had generally higher scores than men, and widowed and separated subjects higher than those who were currently or never married⁴.

Depression was confirmed as a common illness among older people in Europe. A number of other studies have shown poor treatment levels. It was concluded that opportunities for effective treatment were almost certainly being lost.

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Depression in Older Primary Care Patients: Diagnosis and Course

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Depressive symptoms and syndromes in later life are a major public health problem^{1,2}. Primary care clinical settings are especially important venues to better understand depressive psychopathology among older people. Older people with psychiatric disorders utilize mental health services infrequently, especially in comparison with younger persons, yet they are

more likely to see their primary care physicians regularly^{3,4}. Elders who complete suicide have often seen their primary care providers shortly before death, and the majority of them were suffering from depressive conditions at the time of their death⁴. There are also many lines of evidence suggesting that the nature of psychopathology seen in primary care differs from that seen in

EURODEP—Prevalence of Depression in Europe

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The aims of the EURODEP Concerted Action were to use existing studies which had employed the GMS-AGECAT method of diagnosis to assess the prevalence of depression in nine European countries (10 centres)—Liverpool, Amsterdam (sample A), Berlin, Dublin, Iceland, London, Maastricht, Munich, Verona and Zaragoza. Later Tirana was added (not reported here), and five non-AGECAT centres joined after the study commenced—Bordeaux, Oulu, Antwerp, Amsterdam (sample B) and Göteborg.

All subjects were aged 65 or over. In Munich the subjects were aged 85 and above and in Iceland 85–87. The Amsterdam study (sample A) had an upper age limit of 84. All the studies took random samples in the community except Dublin, which sampled a general practice. Sample size varied from 202 in Verona to 5222 in Liverpool, giving a total sample of 13 803. Substantial differences in the prevalence of depression were found, with Iceland having the lowest level at 8.8%, followed by Liverpool, 10.0%; Zaragoza, 10.7%; Dublin, 11.9%; Amsterdam, 12.0%; Berlin, 16.5%; London, 17.3%; Verona, 18.3%; and Munich, 23.6%. When all five AGECAT depression levels, including both subcases of depression and cases, were added together, five high-scoring centres emerged, namely Amsterdam, Berlin, Munich, London and Verona (30.4–37.9%) and four low-scoring centres, Dublin, Iceland, Liverpool and Zaragoza (17.7–21.4%). There was no constant association between prevalence and age. A meta-analysis of the pooled data on the nine European centres yielded an overall prevalence of 12.3% (95% CI, 11.8–12.9); for women, 14.1% (95% CI, 13.5–14.8) and for men, 8.6% (95% CI, 7.9–9.3%)¹.

The proportions of depressive symptoms were found to vary between centres. In Amsterdam, for example, 40% of a general population of older people admitted to depressive mood, compared to only 26% in Zaragoza. Symptoms such as “future bleak”, “hopelessness”, “wish to be dead”, were generally rare, but the last reached higher levels in Berlin, Munich and Verona. Sleep disturbance was admitted by only 15% of the population in

Dublin, but 54% and 60% in Munich and Berlin. Large differences for some depressive symptoms were found within the very old populations, with lower levels in Iceland and higher levels in Munich. Overall, the levels of depressive symptoms among over 60% of the older general population of Europe were low, so that pejorative stereotypes of old age in Europe were not upheld².

In order to include non-AGECAT centres, attempts were made to harmonize the depression measures that had been used with items from the Geriatric Mental State examination. A scale was constructed, the Euro-D scale³. The scale appeared to work well and was applied to data from 21 724 subjects. Euro-D scores tended to increase with increasing age, unlike the levels of prevalence of depression. Women had generally higher scores than men, and widowed and separated subjects higher than those who were currently or never married⁴.

Depression was confirmed as a common illness among older people in Europe. A number of other studies have shown poor treatment levels. It was concluded that opportunities for effective treatment were almost certainly being lost.

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Depression in Older Primary Care Patients: Diagnosis and Course

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Depressive symptoms and syndromes in later life are a major public health problem^{1,2}. Primary care clinical settings are especially important venues to better understand depressive psychopathology among older people. Older people with psychiatric disorders utilize mental health services infrequently, especially in comparison with younger persons, yet they are

more likely to see their primary care physicians regularly^{3,4}. Elders who complete suicide have often seen their primary care providers shortly before death, and the majority of them were suffering from depressive conditions at the time of their death⁴. There are also many lines of evidence suggesting that the nature of psychopathology seen in primary care differs from that seen in

psychiatric or residential care sites^{3,4,6,7}. Thus, to better understand the nature of depression seen among primary care elders, we must study subjects recruited from primary care practices⁸⁻¹¹.

A growing number of studies have examined the prevalence of mood disorders among older primary care patients. Similar to younger and mixed-age primary care samples, major depression is common, with a point prevalence of 5–10% in the elderly^{6,12,13}. An even greater number of patients have a history of major depression but are currently not fully syndromic, i.e. the point prevalence of major depression in partial or full remission is approximately 12%. Thus, one-fifth of all older persons seen in primary care settings have a depressive condition that, at the least, requires vigilance and education regarding recurrence and may require acute, continuation or maintenance therapy.

The prevalence of so-called “lesser” depressive conditions is also considerable among primary care elders. Dysthymic disorder is relatively uncommon, however, with an estimated point prevalence of 1%, of which half is co-morbid with superimposed major depression⁶. This low prevalence primarily reflects that the specific entry criterion of dysthymia seems to be less applicable to elders, where they fail to fulfill the requirement that depressed mood occurs for “most of the day, for more days than not . . . for at least two years”. However, studies using any of various definitions of “minor”, “subsyndromal” or “subthreshold” depression have noted point prevalences comparable to those for major depression^{7,14,15}. For example, one study used the criteria set for minor depressive disorder proposed in the Appendix to DSM-IV, finding a prevalence of 5%⁷. The same group also arbitrarily defined a subsyndromal group comprising patients scoring >10 on the Hamilton Rating Scale for Depression (but *not* meeting criteria for major or minor depression) and noted a point prevalence of 10%. Some have raised concerns that classifying such “lesser” depressive symptoms as mood disorder is bringing “normal” age-related distress under the rubric of psychopathology. In fact, patients with minor and subsyndromal depression suffer both medical co-morbidity and functional disability comparable to that of major depression (and substantially greater than non-depressed controls), suggesting that these are conditions of considerable clinical importance, albeit a heterogeneous and as-yet poorly characterized group.

While studies of younger or mixed-age groups of primary care patients have demonstrated that major depression has high rates of persistence in both remitting–recurring and continuous patterns, broadly comparable to the chronicity seen in psychiatric treatment settings, much less is known about elders. So-called “lesser depressive symptoms” were powerful risk factors for subsequent new onset, or recurrent, diagnosable depression disorders in mixed-age or younger adults^{16,17}. However, few investigations have focused on older persons or disentangled the findings of their older subjects from their younger subjects. Two prospective studies^{18,19} of older depressed subjects found that depressive symptoms (at 33 months and 9 months, respectively) had considerable rates of persistence but also of remission and recurrence. Both studies assessed depression solely by use of self-report depressive symptom scales, and therefore were unable to determine specific depressive diagnoses. These results were also limited by the potentially reduced validity of self-report methodology regarding depressive symptoms among older persons. One recent report¹², using well-operationalized measures to establish a research diagnosis of major depression among older patients seen at university-affiliated internal medicine centers, found that at 6 month follow-up 38% of subjects with major depression continued to be fully syndromic, and only 11.5% were fully recovered. Preliminary data from our group showed that more than half of older primary care patients suffering major, minor or subsyndromal depression still suffered clinically significant depressive symptoms at 1 year (Lyness *et al.*, manuscript in

preparation). In sum, the course of major and other depressions among primary care elders is largely unknown, but available evidence suggests considerable persistence, as well as variability of symptoms, over time.

Similarly, examination of specific predictors of outcome in this group is largely lacking. Medical illness burden is the most powerfully and consistently identified factor associated with the presence or course of depression in later life^{1,20,21}. However, most studies of the relationship of medical illness to depression in later life have used psychiatric patient populations or mixed-community samples, rather than subjects from primary care settings. There have been cross-sectional studies demonstrating an association between medical illness burden and depression in primary care elderly^{6,12,22}. Two published longitudinal investigations of older primary care patients^{18,19} found that medical status (number of illnesses and self-health perception, respectively) predicted depressive symptoms at 33 and 9 month follow-up, respectively. However, these studies were limited by the use of relatively crude proxies for medical co-morbidity and by a lack of diagnostic assessments for depression. Thus, the predictive role of medical illness (measured by validated instruments) with regard to major or subsyndromal depression in primary care elderly remains to be defined. Similarly, other factors that predict depression outcome in younger adults, or in older psychiatric or nursing-home patients, include functional disability, personality trait neuroticism and social support; with few exceptions, these predictors have not been studied in primary care elders.

Little is known about treatment, too. In our above-cited study, less than half of patients with current major depression were prescribed antidepressant treatment⁷, consistent with findings in younger people that depression is frequently undertreated. A wide variety of physician, patient and physician–patient interaction factors may underlie this low treatment rate, but these remain to be defined rigorously through a variety of quantitative and qualitative study methods. Also, antidepressant treatment rates were similar for major, minor and subsyndromal depression groups, despite the lack of empirical support for medication efficacy for the latter conditions. This suggests that the primary care physicians did not discriminate among these diagnostic categories. Rather, they appear to make treatment decisions based on factors that do not directly relate to the symptomatic or syndromic severity of their patients’ depressive conditions.

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Neurochemistry

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As a preface to this chapter, it is important to consider that major depression is characterized and defined by descriptive, not biological, criteria, making it unlikely that a neurochemical finding could adequately characterize the disorder. The underlying validity of the diagnosis is assessed against other descriptive features of the illness such as symptoms, course, treatment outcome and biological changes. Yet people who manifest similar clinical phenomenology may not necessarily share similar pathophysiology. Major depression is heterogeneous in its expression, possessing various phenomenology, family history and course. Therefore, the underlying biology of each subtype would be expected to be distinct from others. This, however, is often not the case.

In the elderly, a neurochemical characteristic of depression would have to be sufficiently specific to distinguish depression from dementia or from secondary depression. In addition, neurochemical differences would be expected between late-onset and early-onset depression, or delusional and non-delusional depression, thus helping to validate these putative subtypes.

PRIMARY DEMENTIA VS. PRIMARY DEPRESSION IN THE ELDERLY

Both of these disorders share certain neurochemical characteristics and behavioural symptoms. For example, noradrenergic deficits are common to both disorders. Interestingly, greater neuronal loss in the nucleus locus coeruleus, the main noradrenergic outflow to the cortex, occurred in Alzheimer's disease (AD) patients who showed clinical manifestations of depression before death than in those who did not^{1,2}. Similarly, primary dementia patients with major depression had over a 10-fold greater reduction in norepinephrine (noradrenaline) than non-depressed dementia patients³, suggesting that, at least among elderly patients, noradrenergic functions may play a role in secondary depression.

Cerebrospinal fluid (CSF) studies demonstrate decreased concentrations of somatostatin in both major depression and AD patients, often with no differences in monoamine metabolites⁴.

SECONDARY VS. PRIMARY DEPRESSION

Significant depressive symptomatology is common in older groups, who are more likely to be medically ill, and depressive symptoms in medically ill patients range from 10% to 50%⁵. However, the neurochemistry associated with depression in the

elderly, such as the dexamethasone suppression test (DST), thyroid releasing hormone (TRH) test and platelet monoamine oxidase (MAO) activity, as discussed below, may change in physical disease as well as in depression, so that their adequate evaluation requires consideration of the effects of medical illness⁶⁻⁸.

LATE-ONSET VS. EARLY-ONSET DEPRESSION

Early-onset depressives, i.e. patients with depression first occurring earlier in life, may be biologically distinguished from similarly-aged patients with late-onset depression. For example, late-onset elderly patients have a lower sedation threshold to amobarbital (both before and after treatment) than early-onset depressives⁹; they may have greater cortical atrophy and ventricular enlargement; and they may have relatively increased platelet MAO activity (see below).

DELUSIONAL VS. NON-DELUSIONAL DEPRESSION

Delusional depression may represent a distinct clinical subgroup and be more common among patients with late-onset depression¹⁰. Yet there is little evidence for biological differences between groups of depressed patients with and without delusions. For example, urinary MGPG (3-methoxy-4-hydroxy-phenylglycol), a metabolite of norepinephrine, has been reported to be reduced¹¹, increased¹² and not different¹³ in delusionally depressed patients. Similarly, serum levels of dopamine- β -hydroxylase have been reported to be both lower¹⁴ and not different¹⁵ in delusional depression.

BIOLOGICAL CHANGES IN AGING AND DEPRESSION

Many of the neurobiological changes associated with aging are similar to those that occur with depression. For example, normal aging and depression are both associated with decreased brain concentrations of serotonin, dopamine, norepinephrine, their metabolites, increased brain MAO-B activity, increased hypothalamic-pituitary-adrenal (HPA) activity and increased sympathetic nervous system activity¹⁶.

HYPOTHALAMIC–PITUITARY–ADRENAL (HPA) AXIS

Depression across the age range is associated with hyperactivity and dysregulation of the HPA axis, characterized, in part, by elevated plasma and urinary cortisol, increased corticotropin-releasing hormone (CRH), blunted corticotropin (ACTH) response to CRH and resistance of cortisol to suppression by dexamethasone. Increased cortisol secretion is probably the most consistently observed physiological abnormality in patients with major depression (see Chapter 81 on dexamethasone suppression test).

Total urinary free cortisol may reliably distinguish depressed from non-depressed individuals in clinical studies of mixed-aged adults. The levels of cortisol, furthermore, appear to correlate with the severity of depression, the presence of psychotic features, and with cognitive impairment¹⁷. Anatomical imaging research shows correlations of cortisol with brain ventricular enlargement¹⁸.

One way in which aging and depression may interact is as follows. It is possible that normal aging is associated with enhanced limbic–hypothalamic–pituitary–adrenal axis activity, possibly due to age-associated neuronal degeneration in the hippocampus. The neuronal degeneration, in turn, may result from increased levels of glucocorticoids associated with both depression and aging. Depressive illness in the elderly may exacerbate this condition, as may repetitive stressful life events.

PLATELET MONOAMINE OXIDASE (MAO) ACTIVITY

MAO catalyses the oxidative deamination of several monoamines and exists in two forms, identified by relative substrate specificity. In the brain, both MAO-A and MAO-B are present; in platelets, only MAO-B is present. In the brain, MAO-B may be more responsible for the degradation of dopamine and MAO-A for norepinephrine.

Platelet MAO activity increases with age, nearly doubles between ages 30 and 80, and is higher in females than males¹⁹. Platelet and brain MAO-B activity is increased in AD, over and above the increase associated with age^{20,21}. Some medical conditions, including liver disease, anaemia, epilepsy and cancer, are also associated with increased MAO activity, whereas in diabetes this may be lower⁷. Although not specific for a particular psychiatric disorder, differences in platelet MAO activity seem to distinguish certain depression populations. Among subgroups of mixed-age, depressed subjects, MAO activity has been reported to be higher in primary depressed, non medically-ill outpatients than in controls, and also higher in secondary than in primary depression^{22,23}. MAO activity may not distinguish endogenous from non-endogenous or delusional from non-delusional depression²², but it may be lower in bipolar compared with unipolar depression²⁴.

Post mortem studies of MAO in the brain have shown that the quantitative distribution of MAO-A in brainstem monoamine nuclei is normal in major depression²⁵. However the density of brain MAO-B in suicide victims showed a positive correlation with age, but was not different between suicides and age-matched controls²⁶.

Elderly female inpatients with primary, unipolar depression, predominately endogenous, whose illness onset was in mid-life, may have lower platelet MAO activity than either later-age-onset depressed females of controls²⁷, suggesting that despite similar clinical presentation, decreased MAO activity may be associated with early-onset depression in the elderly. On the other hand, in elderly depressed female outpatients, MAO activity was higher

than controls, but age of onset was not assessed²⁸. In a depressed population, lower platelet MAO activity predicted the occurrence of neurotic depression (ICD-9) 10 years later²⁹.

In a group of elderly, depressed outpatients, higher platelet MAO activity was correlated with anhedonia, anxiety, a positive family history of depression and response to MAO inhibitors³⁰. Depressed patients with reversible cognitive impairment have higher MAO activity than depressed patients without cognitive impairment³¹. Also, among elderly outpatients, MAO activity was increased in a group with depression secondary to mental illness compared with a primary depression group³². Thus, changes in platelet MAO activity are complex among the elderly depressed population.

Platelet MAO activity may serve as a vulnerability marker for psychopathology in general. Association studies suggested that in the population with age over 40 years, presence of the 165 bp allele of DXS7 at the MAO locus was significantly associated with unipolar depression³³. Both high- and low-MAO activity are associated with increased risk of bipolar disorder, depression and alcoholism in family members.

SEROTONIN

Deficits in the serotonin neurotransmitter system have been implicated both in major depression and in suicide. Numerous studies have demonstrated an association between low platelet 5-HT uptake and depression and a recent study showed higher uptake efficiency in depressed patients with high net uptake rate but similar 5-HT content to normal controls³⁴. Many studies suggest that the serotonin metabolite 5-hydroxyindole acetic acid (5-HIAA) is decreased in the CSF of depressed patients; presynaptic serotonin function is decreased as well³⁵. Specific brain ³H-imipramine binding sites are correlated with serotonin patterns of innervation and are located on presynaptic serotonergic nerve terminals³⁶. The active component for this binding site has a role in modulating neuronal serotonin uptake. Among suicide victims, at least, there is evidence for a decrease in the number of imipramine receptors³⁵, and an increase in the number of postsynaptic serotonin₂ receptors³⁷. This latter finding may represent upregulation due to decreased intrasynaptic serotonin.

The role of 5-HT has also been investigated in genetic, neuroimaging, neuroendocrine and post mortem studies. Serotonin transporter gene polymorphism was not associated with old-age depression³⁸. However, in patients with Parkinson's disease, those with the short allele of the 5-HT transporter promoter region—associated with low 5-HT transporter density—had higher depression and anxiety scores than those without this polymorphism³⁹. Secondary depression to Parkinson's disease, stroke and Huntington's disease in the elderly is associated with low cerebral blood flow and metabolic rate in orbital frontal cortex and basal ganglia, suggesting that, as in primary depression, there is disruption in cortical–basal ganglia–thalamic neuronal loops⁴⁰, abnormalities that are under 5-HT control⁴¹. Poststroke depression is associated with blunted prolactin response to buspirone indicating low 5-HT activity⁴². Serotonin transporter density, measured using paroxetine binding in midbrains of suicides with major depression, were comparable to values obtained in normal controls⁴³.

Although a considerable literature evaluating CSF markers in mixed-age depressed and suicidal patients exists, only one study has focused specifically on elderly depressed subjects. In this study, both CSF 5-HIAA and the dopamine metabolite homovanillic acid (HVA) were lower in elderly depressed patients who attempted suicide than in depressed non-suicidal patients and controls⁴⁴.

PLATELET ³H-IMIPRAMINE BINDING

The ³H-imipramine binding site is found in brain and on platelets, the two sites sharing many characteristics with each other. The binding site is closely related to the presynaptic serotonin-uptake site. Thus, during the last decade, platelet ³H-imipramine binding density (β_{\max}) has been examined extensively as a biological marker for mood disorder. Approximately two-thirds of the studies have reported β_{\max} among certain depressive subtypes^{46,47}, including those with a family history of depression^{47,48}. However, β_{\max} values seem also to be decreased in obsessive-compulsive disorder, anorexia and enuresis. ³H-imipramine binding defined by desmethylimipramine was lower in the putamen of non-violent depressed suicides and those who were antidepressants-treated than normal controls⁴⁹.

The effect of age on platelet binding density has not been systematically investigated. A few studies suggest an increase, decrease, or no change in density with age, but the age ranges have been limited. Animal studies suggest that brain binding sites increase with age³⁶. Elderly unipolar depressed patients also show a 20–42% decrease in binding density when compared to age-appropriate controls, although, again, this is not always so⁵⁰.

Differences among specific depression subtypes in the elderly have not been widely studied; platelet receptor density seems to be somewhat lower in elderly depressed patients compared to a younger depressed cohort⁵¹. Platelet ³H-imipramine β_{\max} may discriminate between depression secondary to medical illness and primary major depression, with density decreased in primary depression patients compared to secondary depression patients and controls³².

It is not known whether this marker represents a trait or a state characteristic. The evidence suggesting state dependence is that depressed patients treated with electroconvulsive therapy showed an increase in binding density⁵². The administration of various medications such as fluoxetine, paroxetine, citalopram and chlorimipramine affect imipramine binding.

Platelet ³H-imipramine binding density does not seem to be affected, overall, in groups of AD patients^{51,53,54} compared to controls, but AD patients with agitation and delusions may have a lower density than AD patients without these symptoms⁵³.

NORADRENERGIC FUNCTION

Studies in depression have failed to demonstrate consistent differences in brain or CSF norepinephrine (NE) or MHPG levels. Normal aging is associated with a decrease in brain NE and a loss of neurons from the noradrenergic nucleus locus coeruleus. In addition, there are decreases in two enzymes required for norepinephrine synthesis, tyrosine hydroxylase and dopa decarboxylase. A post mortem study of depressed patients reported lower NE transporter binding of ³H-nisoxetine in the midcaudal portion of the locus coeruleus (LC) than normal controls, which may be related to low NE availability⁵⁵. Tyrosine hydroxylase (TH) immunoreactivity is reduced in LC in depressed non-suicidal patients indicating low NE availability⁵⁶, whilst its expression is elevated in LC of depressed patients indicating premortem overactivity or deficiency in NE⁵⁷. Neurotrophin 3 which was shown to prevent the death of central NE neurons is low in CSF of elderly patients with major depression⁵⁸.

Platelet α_2 -Adrenergic Binding

Presynaptic α_2 -receptors, in general, seem to function as noradrenergic modulators by regulating NE through a feedback mechanism. Activation of the receptor decreases NE function,

while antagonism increases it. Decreased α_2 -receptor density in brain and platelets may indicate a subsensitive autoreceptor system with increased NE activity, whereas increased α_2 -autoreceptor density may be a characteristic of the putative deficits in central NE function in depression. However, a post mortem study of α_2 -receptor activity in prefrontal cortex and hippocampal cortex in depressed patients showed similar levels to values in normal controls⁵⁹.

α_2 -Adrenergic receptors exist peripherally on platelets, and in most studies, appear increased in depressed patients⁶⁰. Increased platelet α_2 -receptors have also been found in elderly depressed compared to age-appropriate controls⁶¹ (studies that report increased α_2 -receptors tend to use α_2 -agonists in the assay while studies showing no difference tend to use antagonists).

CONCLUSION AND METHODOLOGICAL LIMITATIONS

Limited work has been done in specifically assessing the neurochemistry of depression in the elderly. The biological characteristics discussed here generally discriminate depressed from non-depressed in older as well as in younger groups, but often occur in other disorders as well. Therefore, neurochemistry characterizing depression in the elderly has limited generalizability and validity.

Some of the neurochemical findings described may not be specific to affective disorder in the elderly, but indicate traits that are over represented in groups of depressed patients when compared to the general population. For example, decreased platelet imipramine binding density may characterize people with relative serotonin system deficits, who may be at increased risk for depression, or who have symptoms related to depression. Or, increased platelet α_2 -receptor binding may identify individuals who possess noradrenergic or arousal defects associated with depression.

Whereas platelet, plasma and CSF are sources of assayable human tissue, the relationship of their biochemistry is not well understood. The similar embryological origin of megakaryocytes and neurons has been used to justify or explain platelet research findings. However, these cells and their platelets have differentiated considerably from neurons and are exposed to significantly different physiological influences. Thus, it could be considered remarkable that it is possible to use platelets at all in the manner discussed here.

The majority of these studies are based on cross-sectional assessments during an acute phase of illness, or at death. Longitudinal studies are needed to assess test–retest reliability, state vs. trait and other characteristics. An adequate understanding of the neurochemistry of depression in the elderly must address age effects and the effects of depression subtypes, and must adequately explain how the neurochemistry is related to the depression.

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Neuro-imaging

Neuro-imaging Studies of Depression

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The use of neuro-imaging techniques in the study of depression has provided a major advance in the elucidation of its functional neuro-anatomy, metabolic correlates and biochemistry. Structural neuro-imaging techniques, such as computed axial tomography (CT) and magnetic resonance imaging (MRI) have demonstrated an association between structural brain abnormalities and depression in patients with unipolar and bipolar disorder¹. Whilst these abnormalities are non-specific, there is evidence that volume reduction in the caudate² and frontal lobe³ are specific abnormalities in depression, particularly in late-life depression. MRI studies have consistently shown an increase in the number and/or severity of signal hyperintensities in the white matter in both unipolar and bipolar disorder⁴. Deep white matter lesions have been shown in elderly patients with unipolar depression predominantly in the frontal lobes and basal ganglia, supporting the notion of frontostriatal dysfunction in depression. These lesions have been shown to be associated with poor long-term outcome⁵. A recent population-based study of over 1000 elderly people showed that those with severe periventricular white matter lesions were three to five times more likely to have depressive symptoms than those with only mild or no white matter lesions, whilst those with severe subcortical but not periventricular white matter lesions were more likely to have had a history of late-onset depression after the age of 60 years than those with only mild or no white matter lesions⁶. Severe white matter lesions are thought to represent vascular abnormalities, findings that support the notion that vascular pathology contributes to the aetiology of late-life depression.

Functional neuro-imaging techniques have also advanced our knowledge of the pathophysiology and chemical pathology of depression with the introduction of single-photon emission computed tomography (SPECT), positron emission tomography (PET) and magnetic resonance spectroscopy (MRS). Several studies have shown an association between depression, low cerebral blood flow and low metabolic activity. An extensive review of the literature showed evidence for low cerebral activity in the frontal lobes in unipolar and bipolar depressive patients, with an inverse correlation with increased severity of depressive symptoms and an association with low activity in basal ganglia and temporal and limbic regions⁷. Low neurostriatal cerebral blood flow was particularly associated with psychomotor slowing⁸ and an association between hypofrontality and negative symptoms in depressive patients⁹. This abnormality is of special interest in the elderly because of the interface between depression and dementia in this age group.

We compared the cerebral perfusion in a group of elderly depressed patients with that in age matched healthy subjects and

Alzheimer's disease (AD) patients, using SPECT methodology and hexamethyl propleamine oxime (HMPAO) as the radioligand¹⁰. The regional cerebral perfusion values of depressed patients were intermediate between those of the healthy control subjects and the AD patients. This is consistent with the intermediate position of the depressed group on other measures, e.g. radioattenuation on CT scan. In addition to the global impairment in cerebral perfusion, there are topographical abnormalities but the cerebral region thus affected is seen to vary in different studies. Sackeim *et al.*¹¹ have argued that the traditional statistical paradigms, concentrating on identifying specific brain regions with higher or lower perfusion compared to control subjects, have failed to examine the important issue of identifying the abnormal patterning of regional activity, which would reflect the activity of functional neural networks. Using a novel scaled subprofile model based on factor analytic technique, they demonstrated topographical abnormalities in the temporoparietal area, largely consisting of polymodal association cortex with strong reciprocal connection with the prefrontal polymodal association cortex. The study by Baxter *et al.*¹² on glucose metabolism using PET strongly suggests that the biological substrate common to depressive states in different patient groups is a reduction in glucose metabolism in the left prefrontal cortex.

The impairment in cerebral perfusion correlates positively with the endogenicity score but not necessarily with the severity of depressive illness¹³. This is in contrast with studies on cerebral glucose metabolism using PET, reporting a positive correlation between the severity of depressive symptoms and glucose metabolic rate. There are some reports on abnormality in the anteroposterior gradient in cerebral perfusion in depressive illness, but these are not consistent.

The temporal sequence of the perfusion abnormality and clinical depression is yet unclear. The impairment in cerebral perfusion improved with clinical recovery, suggesting that the continuing impairment can predict chronicity of the course or frequent relapse of depression. This notion can be examined in further longitudinal studies.

SPECT and PET techniques have also been utilized to investigate the chemical pathology of dopamine and serotonin systems in depression. SPECT studies of D2 receptors showed increased D2 receptor density in the striatum, reflecting reduced dopamine function¹⁴, and an association between increased D2 receptor binding in the left striatum in the anterior cingulate gyrus and clinical recovery with selective serotonin reuptake inhibitors¹⁵. Studies of the dopamine transport receptors in depression showed increased receptor density in the basal ganglia¹⁶. Studies

of the serotonin system showed a number of interesting associations. A SPECT study of the brain serotonin transport receptor reported reduced density in depression¹⁷, whilst a PET study of the serotonin type 2 receptor function showed a decrease in receptor density following treatment with desipramine, particularly in the frontal region¹⁸. Studies of the 5-HT_{1A} system showed a reduction in receptor density in the mesiotemporal cortex in bipolar patients, and in unipolar patients with bipolar relatives¹⁹ and an association between widespread reduction of 5-HT_{1A} receptor binding and depression in medicated and non-medicated depressed patients²⁰. Finally, studies of MRS depicting biochemical changes have detected an association between depression and reduced glutamine and glutamate, using proton-MRS²¹, and increased phosphomonoesters and decreased ATP values in the frontal lobes of patients with major depression using ³¹P-MRS²².

In summary, the findings of neuro-imaging studies in depression add credence and support to the biological basis of depression, refine its diagnostic subgroups and inform its treatment with more specific pharmacological treatments.

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Is Imaging Justified in the Investigation of Older People?

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Dementia is still diagnosable only by clinical examination, but a further battery of diagnostic tests may then be required to clarify the aetiology and identify potentially treatable cases. Any assessment of the cost-effectiveness of these further tests, including neuro-imaging, must compare the relative value of routine testing in all patients against selective testing in cases where there is a high index of suspicion or diagnostic doubt.

Routine testing will detect the small number of patients (about 1%) with a reversible cause for their dementia, but is burdensome, especially in the elderly, and may raise false expectations and lead to false-positive results. It would also identify early cases suitable

for drug therapy. Selective testing will result in some treatable cases being missed, but causes less general risk and discomfort and may be more cost-effective, especially in those countries where neuro-imaging remains expensive and difficult, or even impossible, to access^{1,2}.

Providing cost-effective and accurate diagnosis of dementia therefore presents the clinician with a dilemma. Can neuro-imaging be included in a battery of ancillary investigations devised for routine use in all cases of dementia, or need such tests be carried out only as clinically indicated? The fact that definitive data on the sensitivity, specificity and cost-effectiveness of various

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neuro-imaging techniques are still needed³ only adds to the problem.

The advent of treatments for dementia and the consequent need to identify early and potentially treatable cases has shifted the assessment of cost-effectiveness from investigation to treatment. The cost-benefit assessment of early treatment of dementia to a health economy has yet to be fully assessed but is likely to far exceed that of any neuro-imaging screening programme. A recent review⁴ neatly encapsulates what is now as much a socioeconomic as a medical dilemma in a single statement: "How much is society willing to spend?"

Given that these socioeconomic drivers are unlikely to change, clinicians are now seeking to devise assessment protocols for dementia that best address the conflict. Unfortunately, we are still some distance from achieving a consensus view. Some authors⁴ advocate routine neuro-imaging as part of a screening procedure in all patients in order to "include in" all treatable cases, perhaps avoiding litigation. Others, when attempting to define practice parameters, have regarded neuro-imaging as optional in the differential diagnosis of dementia⁵. Subsequent analysis showed that imaging studies did improve diagnostic accuracy but only with a significant increase in cost⁶.

Considerations of cost and, in less well-developed countries, of access continue to determine whether neuro-imaging in dementia is used as a universal screen or only when clinically indicated. Various protocols indicating that neuro-imaging is not required in all patients, but should be a first-line test, especially in younger patients and those cases where there is diagnostic doubt on clinical grounds, have been suggested⁷⁻¹⁰. It is likely that, as costs of treatment increase relative to those of investigation, the use of

neuro-imaging as a screening tool rather than a diagnostic aid in dementia will increase.

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Clinical Features of Depression and Dysthymia

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Depression *per se* is not characteristic of later life and should not be interpreted as “normal”. Most normal adults make a satisfactory adjustment to the fact of growing older; after all, it is preferable to the alternative. Depression in old age may represent a continuation of a process that began in early life. For example, dysthymia as a chronic condition may extend into later life, or a continuation of cyclical changes of mood may be experienced, with periodic exacerbations of unipolar or bipolar depression. Additionally, some types of depression may originate in later life and the symptomatic presentation may or may not vary from the classical symptomatology. Depression and dysthymia may arise as a component of other psychiatric disturbances. Misdiagnosis or lack of referral for treatment of depression and dysthymia in older adults is due both to the context in which depression appears and the manner of presentation of symptoms¹. Depression is a spectrum of disorders, rather than a unitary construct, and multiple biopsychosocial factors will influence how clinical features are manifested, as well as response to treatment. A depressive disorder may be diagnosed in accordance with current clinical criteria only if the depressive symptoms actually interfere with work, family or social functioning, or if the individual seeks professional treatment or has received medication for the disturbance².

The existing nomenclature may not accurately reflect depressive syndromes in late life. A number of well-established clinical syndromes are documented in the literature, for example depressive pseudodementia, somatization with underlying depression, and other forms of masked depression in individuals who deny depressive symptomatology. Subsyndromal depression may also occur in old age and can progress to a full syndrome of depression. Because of these potential differences in the presentation of depression in late life, epidemiologic data suggest that the elderly have a lower prevalence of major depression than do younger populations¹⁻⁴. The literature has also consistently reported that a low proportion of clinically depressed elderly individuals seek or are referred for psychiatric care⁶. However, despite the inherent problems of describing depression and dysthymia in late life, and the presumed low rates of recognition or referral, depression is probably the most common psychiatric disorder in late life⁷⁻¹⁰.

The clinical features of various forms of depression will be described using an outline form. The three clinical entities relevant to the current nomenclature and older adults are: *major depression*, unipolar or bipolar, single episode or recurrent, with

or without melancholia, with or without psychotic features; *dysthymic disorders* or depressive neurosis; and *other atypical forms of depression*, i.e. somatization, cognitive impairment, including the reversible syndrome of dementia (pseudodementia) mood syndromes secondary to a general medical condition, and subsyndromal depression. The fact remains that in clinical practice considerable overlap exists for these syndromes, boundaries are often not distinct, and a spectrum of symptomatology is observed.

CLINICAL FEATURES OF THE DEPRESSIVE SPECTRUM

Symptoms of Grief and Pathological Bereavement

The hypothesis that demoralization and despair result from losses or incapacities due to aging seems to have no basis in empirical data. In fact, most surveys of older adults show older individuals to be more contented with their life situations than are those in earlier stages of the life cycle⁹⁻¹¹. Perhaps when life events are anticipated and rehearsed, grief work can be completed before the loss with effective coping strategies. Grief, a psychologic response to the loss of a loved one, of possessions or of health itself, is associated with feelings of sadness, transient symptoms of gastrointestinal complaints, weight loss and difficulty in sleeping. Symptoms are often intense for several weeks and then typically begin to improve¹². Many losses can lead to fear, demoralization or loneliness. In vulnerable individuals they may result in the onset, worsening or persistence of a mood disturbance. Pathological grief may include overactivity without a sense of loss, acquisition of symptoms belonging to the last illness of the deceased, frank psychosomatic illness, an alteration of relationships with family and friends with hostility towards specific individuals, and a persistent loss of patterns of social interactions¹³⁻¹⁵.

Because older persons themselves, their families or even healthcare professionals can understand the difficulties experienced, depression occurring in the wake of such significant loss may be considered as “normal”. This uninformed perspective works against making the diagnosis of clinical depression and interferes with the effective treatment of this painful and potentially life-threatening condition. Spousal bereavement is a common occurrence of late life, associated with prolonged personal suffering, declining mental and physical health and

increased risk of mortality¹⁶. Among elderly widows and widowers, 20% meet the criteria for major depressive syndrome 2 months after their loss, and one-third of these individuals develop persistent depression for a year or longer¹⁷. Generally, individuals who are at greatest risk for persistent depression have worse health, more functional and social difficulties and more protracted grief than do bereaved individuals who are not depressed. Whenever major depression occurs, even if precipitated by a significant loss, it should be considered an illness and treated accordingly. As a rule of thumb, if a depressive syndrome occurs within the first 2 months after bereavement and lasts less than 2 months, it should be considered part of the normal grief process and not aggressively treated. However, even within the first 2 months, the depression should be considered a clinically important, treatable syndrome if it is severe or, associated with psychomotor retardation, is accompanied by feelings of worthlessness or suicidal ideation, or occurs in a person with a history of previous major depressive episode⁵. Persistence of severe symptoms beyond several months suggests a condition more severe than the normal grief process.

Symptoms Reflecting Adjustment

Disorders with Depressive Symptoms

The role of social or situational stressors contributing to depression in late life is not well established¹⁸. Situational stressors may include limited mobility, sensory deprivation, retirement, economic constraint, changing or unsatisfactory living conditions, social isolation, marital difficulty, loss of significant loved ones and/or rejection by children¹⁹.

Of much greater frequency, however, is the development of depressive symptomatology secondary to general medical conditions. When the depressive symptoms accompanying physical illness dramatically exceed the level of symptoms expected, then the diagnosis of adjustment disorder is appropriate. General medical illness resulting in functional disabilities is believed to represent the more significant stressor, contributing to the loss of resources necessary for the maintenance of self-esteem^{11,20}. Essentially, depression interferes with basic abilities to think, eat, sleep, love, interact with others, maintain a sense of purpose, experience gratification and maintain self-responsibility²¹. Consistent with the medical outcomes studies²², depressive symptoms are associated with much social and physical dysfunction, days spent in bed and even physical pain as adverse as any general medical condition. Co-morbid medical illness increases the chronicity and refractoriness to treatment and it may slow recovery rates for patients with various conditions, including heart disease, stroke, hip fracture and dementia⁵.

The resolution of stressful events probably depends on a variety of factors, including genetic predisposition, prevailing early life experiences, adequacy of previous adaptive and coping mechanisms, patterns and profiles of premorbid personality and the presence or absence of the support system. Not only may family or social dysfunction influence whether depressive symptoms are experienced by an older adult, but family and social support is apparently critical to the successful outcome in treating the depressed elderly. Blazer has outlined six critical areas as follows: (a) the presence of members of the family or social network who will be available; (b) the interaction of the older adult with family members and other individuals within their community; (c) the integrity of the overall family system and social system; (d) the family and social values regarding the psychiatric disorder; (e) the degree of family and social support with respect to tolerance of symptoms; and (f) other coincidental stressors or life events encountered by the family and social system²³.

Symptoms of Dysthymia

Dysthymic disorder is believed to be less frequent in older adulthood³. Co-morbid general medical conditions, cognitive disorders and frequent adverse life events, e.g. bereavement, make the diagnosis of dysthymic disorder quite difficult. Some associated features may include the presence of major chronic stressors, increased physical impairment and more symptoms of anxiety²⁴. Generally, dysthymia is less associated with co-morbid Axis I or co-morbid Axis II disorders, and patients with dysthymia tend to be seen in primary care settings rather than by psychiatric specialists²⁵. The usual psychologic mechanisms of dysthymia, i.e. self-reproach, guilt and the turning inward of hostile feelings towards loss, are not prominent in later life. However, psychodynamic explanations, i.e. loss of self-esteem and the inability to defend oneself against threats to security, are important in the pathogenesis of late-life dysthymia²⁶. The role of narcissistic pathology in the etiology of late-life depression may also contribute significantly to episodes of recurrent depression or defensive grandiosity in response to minor disappointments, or due to self-consciousness or overdependence on approval from others for maintenance of self-esteem, and as a consequence of transitory periods of fragmentation and discohesiveness of the self²⁷. Cultural or developmental factors may also contribute to dysthymia. For example, the development of habit patterns emphasizing activity, productivity and achievements can lead to depression and despair at retirement or with the cessation of parenting²⁸. Also, as opportunities to interact with cohorts decline, an older individual may not reconcile generative disappointments, and may not proceed from Erikson's²⁹ developmental stage of generativity to that of integrity (vs. despair)²⁹.

Symptoms of Major Depression

Major depression is characterized by a persistently depressed mood, loss of interest in usual activities, guilt or other pessimistic thoughts, with accompanying vegetative symptoms reflecting preoccupation with somatic complaints, persistent sleep disturbance, decreased appetite with weight loss, decreased libido, anhedonia and inability to concentrate. Psychotic and delusional symptoms have been reportedly observed with greater frequency in major depressions occurring in later life⁴. Delusional themes are frequently either somatic or persecutory, and are less often characterized as delusions of guilt, sin, poverty, nihilism or jealousy³⁰. Feelings of guilt, obsessive rumination, agitation and ideas of reference are frequently found in older adults with major depression. Perceptual disturbances, e.g. hallucinations, may be present but are less frequent than other psychotic symptoms. The prevalence of major depression in the oldest-old is lower than for persons in mid-life, but may be somewhat higher for those who are "young-old", usually estimated at 2–5%^{31,32}. Patients with major depression are thought to be more likely to exhibit suicidal ideation. Suicidal ideation is strongly associated with completed suicide. One study has suggested that correlates of suicidal ideation include psychomotor retardation, a history of dysthymia, a previous psychiatric inpatient stay, being a "younger-elder" and having symptoms that reflect feeling "guilty, sinful or worthless"³³. These correlates of suicidal ideation may be present among individuals who have dysthymia and other minor forms of depression as well.

Symptoms of Bipolar Affective Disorder

Bipolar disorder may be more common in the elderly population than currently recognized. Symptoms of the depressed bipolar

elderly patient may include profound psychomotor retardation, even to the point of frank stupor. This psychomotor retardation mimics both dementia and the inanition associated with physical disease. Mood can be elevated or irritable but may be rather labile, showing a picture of depressive admixture^{34,35}. These depressed patients may be unable to provide a meaningful history and/or cooperate with diagnostic procedures. Similarly, the elderly, manic patient may make inappropriate sexual comments and/or advances, or be agitated and/or assaultive, and as a result be regarded as cognitively impaired. Although bipolar patients have usually had a history of mood disturbances in the past, these episodes may have been “forgotten” or repressed by both the patients and primary relatives³⁶.

LATE-ONSET DEPRESSIVE ILLNESS

Depression in late life may be characterized as of early or late onset. Early-onset depression which recurs in later life may have symptoms similar to previous episodes. Differences in the clinical presentation may be observed in both early-onset and late-onset depressives³⁷. Genetic predisposition, significant losses or multiple life stressors may interact with age-related biological vulnerabilities. Biological changes due to the physiologic effects of aging have been verified by the study by Schneider³⁸, showing that unmedicated, elderly depressed subjects demonstrated higher monoamine oxidase (MAO) activity than sex- and age-comparable controls.

Recent studies have proposed that neurobiologic and/or psychosocial factors may predispose an older individual to depression or dysthymia³⁹. MRI-defined vascular depression has been identified as a late-onset, non-psychotic type of depression, seen more often in individuals with no family history of depression, together with symptoms of anhedonia and increased psychosocial impairment⁴⁰. Other studies have verified the increasing incidence of depression among individuals with cerebrovascular risk factors, MRI findings associated with vascular disease and symptoms of apathy, together with diminished life quality⁴¹. Central nervous system degeneration, for example, from the biochemical changes in Alzheimer’s disease or other complaints may predispose to an increased incidence of depression⁴².

Age of onset has been used as a correlate for late-life depressive symptomatology. Depressive symptoms may be found to be more frequent among the old-old compared to the young-old, with approximately 20% compared to less than 10% in the community⁴³. This higher frequency of depression among the old-old may be explained by a higher proportion of women, more general medical problems, more cognitive impairment and lower socioeconomic status. When these factors are controlled, no relationship exists between depressive symptoms and advancing age⁴⁴. Nonetheless, depression is associated with disability among the old-old and a number of studies have illustrated the association between depression and frailty, functional disability and co-morbid general medical problems and/or cognitive impairment⁴⁵⁻⁴⁷. Essentially, having more than two previous episodes as compared to two or less is related to younger age, earlier age of onset, dysthymia, feelings of worthlessness, difficulty concentrating, slowed thoughts, suicidal ideation, symptoms of anxiety and decreased perception of social support. Patients with multiple recurrent episodes are also thought to be at higher risk for more severe illness⁴⁸. Late-onset depression is more frequently associated with structural brain changes and cerebrovascular disease, while early-onset depression seems to be more influenced by family and genetic factors. Compared with early-onset depressives, patients with late-onset depression tend to show more loss of interest, less pathological guilt, more psychosis and more generalized

Table 74.1 Somatization: principles of clinical management

1. The presentation is considered in the context of psychosocial factors, both current and past.
2. The diagnostic procedures and therapeutic interventions are based on objective findings.
3. A therapeutic alliance is fostered and maintained involving the primary care and/or psychiatric physician.
4. The social support system and relevant life quality domains* are carefully reviewed during each patient contact.
5. A regular appointment schedule is maintained for outpatients, irrespective of clinical course.
6. The patient dialogue and examination and the assessment of new symptoms or signs are engaged judiciously, and usually primarily address somatic rather than psychologic concerns.
7. The need for psychiatric referral is recognized early, especially for cases involving chronic symptoms, severe psychosocial consequences or morbid types of illness behavior.
8. Any associated, coexisting or underlying psychiatric disturbance is assiduously evaluated and steadfastly treated.
9. The significance of personality features, addictive potential and self-destructive risk is determined and addressed.
10. The patient’s case is redefined in such a way that management, rather than cure, is the goal of treatment.

* Quality of life is an elusive concept but includes the psychosocial domains of occupation, leisure, family, marital, health, sexual and psychological functioning. From Folks *et al.*⁵⁵, with permission.

anxiety. These correlates of depressive symptomatology based on age of onset suggest a certain heterogeneity in depression of old age.

Atypical Forms of Depression

Depressive illness, particularly of late onset, may present without prominent mood disturbance¹⁹. Atypical forms of depression, or masked depression, are thus common among older individuals¹⁹. Masked depression is characterized by the denial of feelings of depression or the lack of complaints of sadness or dysphoria. The dysphoric affect is often prominently masked by somatic complaints, e.g. fatigue, pain, gastrointestinal upset, concentration difficulties or diminished energy⁴⁹. These individuals who manifest prominent somatic symptoms of depression are prone to attribute symptoms of depression to their medical illnesses¹⁰. Two forms of somatization, hypochondriasis and conversion, may predominate in the clinical picture. The clinical approach to somatization is outlined in Table 74.1.

Hypochondriasis is a common form of masked depression in the elderly and may increase the risk for attempted suicide⁵⁰. This “secondary” form of hypochondriasis must be differentiated from primary hypochondriasis, a persistent somatoform disorder that tends to have its onset in the third or fourth decade of life and persists. Hypochondriasis *per se* is *not* more prevalent in the elderly and its onset in later life should not be considered a part of normal aging, but rather reflective of psychologic distress, particularly depression⁵¹⁻⁵³. Conversion symptoms or pain that occurs in an elderly person should also raise the question as to the presence of underlying depression, even when no prominent mood disturbance is found^{54,55}. In the nursing home setting, a patient may not meet the clinical criteria for depression on patient interview but be observed to have depressive symptoms in the context of somatic complaints. Apathy, withdrawal and isolation may be clues that depression is present. On the other hand, a patient in a long-term care facility may become abruptly agitated with sleep disturbance and prominent somatic complaints, which should increase an index of suspicion for depression.

Table 74.2 Diagnostic features distinguishing depression from dementia

Depression	Dementia
Depressive symptoms	Euthymia
Subacute onset	Insidious onset
History of depression more	History of depression less
Aphasia, apraxia, agnosia absent	Aphasia, apraxia, agnosia present
Orientation intact	Orientation impaired
Concentration impaired	Recent memory impaired
Patient emphasis on memory complaint	Patient minimizes memory complaint
Patient gives up on testing	Patient makes effort on testing

Adapted from Wells⁵⁶.

Depression in late life may be masked not only by somatic complaints but also by cognitive difficulties, reflecting yet another atypical form of depression observed in older adults. Difficulty in concentrating or memory loss may result in the clinical picture of “depressive pseudodementia”⁵⁶. This *dementia syndrome of depression* represents a reversible syndrome of dementia that may be clinically indistinguishable from irreversible dementias (Table 74.2)^{57,58}. These individuals frequently make little effort to cooperate with mental status examination and the patient answers “Don’t know” to many questions. However, notable losses of both recent and remote memory are observed, and these patients may show marked variability on the performance of tasks of similar difficulty. In some cases, depression may be diagnosed retrospectively after a favorable response to a trial of antidepressant medication¹⁹. Further complicating the issue, in about 20% of cases of true dementia a major depression may coexist⁵⁹. In these cases, both mood and function may improve when treated with an antidepressant therapy, but the basic cognitive impairment remains.

Depression Secondary to a General Medical Condition

Depression may be intimately related to general medical disease or other “organic” influences. Of course, the relationship between somatic symptoms in the medically ill elderly and complaints of depression may be complex; older individuals are presumably more vulnerable to the stresses of poor health and disability that interfere with body image, self-esteem and autonomy. Comorbidity is the rule, rather than the exception, and linked significantly to functional decline across multiple parameters with increasing age. Co-morbidity of depression with medical illness is associated with poor physical, mental and social functioning, all of which compound the patient’s ability to enjoy the quality of life. Not only may medical problems act as precipitators of depression, but many direct effects of medical conditions induce a secondary form of depression (Table 74.3)^{9,10}. Medications, including anxiolytics, antihypertensives and neuroleptics, may induce a syndrome of depression. Polypharmacy and drug interaction may further serve to culminate in a depressive illness. The association between depression and general medical–surgical problems may be summarized as follows: (a) physical disease is associated with narcissistic injury, loss of autonomy, pain and fear of impending death, frailty (to be discussed) and diminished quality of life; (b) physical illness may directly (e.g. a cerebrovascular accident) or indirectly (e.g. hypercalcemia) cause a depressive syndrome; or (c) medications used to treat medical diseases may themselves induce depression. These parameters are more significantly affected than in an individual with depression alone or general medical illness without depression⁶⁰. Thus, among individuals with depression in the context of dementia, individuals who experience co-morbid depression and general

Table 74.3 Medications reportedly associated with depression

Cardiovascular drugs	Hormones	Psychotropics
α-Methyldopa	Conjugal estrogens	Analgesics
Reserpine	ACTH (corticotropin) and glucocorticoids	Anti-parkinsonian
Propranolol		Antihistamines
Guanethidine	Anabolic steroids	Benzodiazepines
Clonidine		Other sedative hypnotics
Thiazide diuretics		Typical neuroleptics
Digitalis		
Anticancer agents	Anti-inflammatory/anti-infective agents	Others
Cycloserine	Non-steroidal anti-inflammatory agents	Cocaine (withdrawal)
	Ethambutol	Amphetamines (withdrawal)
	Disulfiram	L-dopa
	Sulfonamides	Cimetidine
	Baclofen	Ranitidine
	Metoclopramide	

Adapted from Agency for Health Care Policy and Research⁷⁸.

medical illness are less active, with fewer social contacts that explain increased disability risk⁴⁵. Physical immobility and social isolation in turn increases disability over time, which further decreases mobility and social interactions, placing the person at risk for physical, psychological and social impairment^{61,62}. For some individuals, the end result is a downward spiral in health, functioning and quality of life, a syndrome described by geriatricians as “frailty and failure to thrive”. This manifestation is characterized by weight loss, weakness, fatigue, inactivity, decreased food intake and depression. Physical signs that may accompany these symptoms include sarcopenia, balance and gait abnormalities, deconditioning and decreased bone mass^{63,64}. Failure to thrive specifies an end-stage of frailty that is characterized by unchecked weight loss, severe muscle wasting, apathetic depression and a host of physiologic abnormalities, including hypoalbuminemia, low creatinine, anemia, bicuspid ulcers and untimely death⁶⁵. Indeed, depression in the elderly is closely related to the state of one’s physical health.

Subsyndromal Depression

Subsyndromal depression is defined as depressive symptoms that do not qualify for a formal mood disorder using current clinical criteria. However, several studies have revealed that subthreshold depressive states can be associated with adverse clinical outcome. Additionally, subsyndromal depression may be a risk factor for subsequent major depression⁶⁶. Subthreshold depressive disorder tends to be a heterogeneous group of milder forms of depression with symptom patterns qualitatively distinct from more severe depressions such as major depression. According to the Berlin aging study, a subthreshold depression can be characterized in two ways among the elderly⁶⁷; first, as a quantitatively minor variant of depression or a depression-like state, with fewer symptoms or with less continuity; second, as a condition qualitatively different from major depression, with fewer suicidal thoughts or feelings of guilt or worthlessness, while worries about health and weariness of living occur with a similar frequency.

CLINICAL FEATURES AND PROGNOSTIC FACTORS

The paucity of controlled studies in the diagnosis and phenomenology of depression and dysthymia in older adults restricts

meaningful prognostication. Depression may be reflected in behavioral changes in the elderly, for example increased clinging, dependent behavior, phobias or avoidance of previously enjoyable activities, perhaps representing a variant of anhedonia. No conclusive evidence has shown that major depression is more chronic in later life; in fact, most psychogeriatricians are convinced that the outcome of adequately treated depression, uncomplicated by medical disease, is as good as the outcome from younger individuals. A caveat is that the risk of recurrence may be greater for older depressives^{68,69}. Blazer²⁶ has noted the tendency for some older individuals to show only partial improvement not fully recovering from an episode of major depression. Thus, in some cases residual symptoms may persist even when response to treatment is definitive—this treatment resistance is more attributed to co-morbidity than to age *per se*.

Accurate and early diagnosis and adequate and aggressive treatment are important considerations in late-life depression and dysthymia. Patients should be screened, with history, questionnaires, observation, collateral information and the use of screening devices, such as the Geriatric Depression Scale⁷⁰ or the Center for Epidemiological Studies Depression Scale (CESD)⁷¹, both of which are validated in older depressives and useful in the overall assessment process. These self-assessment instruments, together with the Hamilton Depression Rating Scale⁷², an interviewer assessment agent, are useful for assignment of psychiatric diagnosis of depression. Other questions, designed to establish cognitive, nutritional and functional status, social function and general health perceptions, together with review of medications, medical work-up and laboratory assessment as indicated, are necessary in early intervention. Keller *et al.*⁷³ reported that depressed patients who did not receive treatment until long after the onset of depressive symptoms had the worst prognosis. These investigators suggested that early intervention is indeed the most effective means of reducing chronicity. Therapy must proceed across multiple domains simultaneously. This includes the mobilization of the social support system, the use of antidepressant medications, psychotherapy and efforts to improve physical or social functioning^{74,75}. Among psychotherapeutic options, cognitive-behavioral therapy has been utilized most, with a few reports involving interpersonal psychotherapy. Other behavioral interventions range from prescribing regular group activities, physical exercise such as walking, and training exercises for increased independent functioning. Although few in number, studies of comprehensive interventions using all of the approaches outlined above have demonstrated efficacy comparable to that found in younger adults^{76,77}. These existing studies are preliminary but promising, in view of the reduction in functional disability and symptomatic relief, together with improved quality of life and, ultimately, reduction in the healthcare cost associated with late-life depression.

CONCLUSION

The categorical approach to depression and dysthymia in late life continues to reflect the current nomenclature. Ultimately, the diagnosis and management of depression and dysthymia is allied with the traditional medical model. This approach enables the clinician to provide specific therapies for distinct diagnostic entities and provides the patient with a number of excellent biological and psychosocial interventions. However, classical symptoms may be overshadowed or replaced by a variety of other symptoms, in which the mood disturbance is not prominent. Thus, the key to the diagnosis and effective intervention of depression in old age is to maintain a high index of suspicion and attend to the functional disability and quality of life issues posed by late-life depression.

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Outcome of Depressive Disorders: Findings of a Longitudinal Study in the UK

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A high prevalence of depressive disorders among the elderly has been reported by different community studies (*see reviews*^{1,2}). More recent epidemiological studies³⁻⁹ using standardized methods have also reported similar prevalence rates of depressive disorders in the range 8–15% in randomly selected community samples. It is not clear whether the high prevalence rate is due to chronicity of the disorder or to a high incidence rate of depression in this age group.

So far there have been few large-scale longitudinal studies of community-based elderly depressed subjects. Most of these studies have reported that 30–60% of the depressed elderly were continually ill¹⁰⁻¹⁵. Some studies have examined the prognostic factors associated with the poor outcome. Physical illness, handicap, female sex, poor social support, high depression score and anxiety and neuroticism, lack of satisfaction with life and feelings of loneliness are some of the factors reported to be associated with the poor outcome^{12,14,16-21}. Kivela²², in her 5 year follow-up study, reported organic outcome in 12% for major depression and 9% for dysthymic disorder. In contrast, Henderson *et al.*¹⁶ found none of their 24 community cases of depression (ICD 10) to be organic cases after 3–4 years. Many of the studies have also reported that the majority of the depressed elderly never receive appropriate treatment^{10,13,23-25}.

Community studies^{15,16,22,26,27} have reported increased mortality in depression. There have been a few relatively shorter (1–3.5 years) follow-up studies^{10,29} which found no significant association between depression and mortality. In another study, when factors such as age, physical health and social conditions were taken into account, depression did not predict increased mortality¹⁸.

Tannock and Katona³⁰ reviewed the literature and concluded that subsyndromal depression is common among the elderly and that very little is known about its nature and outcome.

Where the symptoms of an illness appear to lie on a continuum of severity with normal behaviour, as in depressive illness, variations in disease level might be found between studies because they use different cut-off points and/or record different types of symptoms, rather than because there are true differences in illness levels. Such spurious differences may also arise between different waves of interviewing in the same longitudinal study if the interpretation of symptoms is not standardized, and may also occur if the interviewers of subsequent waves are not blind to the findings of previous waves.

A further problem with such studies is that some psychiatrists assume depression to be a unitary disease but nevertheless do not agree on what conditions are to be excluded; e.g. some would include bereavement and adjustment disorder if the principal symptom is one of depression. Others would distinguish between major depressive disorders and dysthymia, or between endogenous and reactive, or neurotic and psychotic depression and might not recognize brief recurrent depression.

Cole and Bellovance³¹, in their review of five community-based studies of depression in old age, highlighted such methodological limitations. They recommended that a large number of depressed elderly (based on explicit diagnostic criteria and reliable measures) should be followed up for a longer duration with specific outcome

categories. Outcome assessments should be blind and reported at specific time points (e.g. every 6 months) during the follow-up interval. A continuously depressed elderly cohort should be examined closely to determine the sustaining factors of their depression.

This special article contains findings of the outcome data of the depressed elderly in the Liverpool longitudinal study.

The subsample for this study included 120 index AGE-CAT cases of depression and 47 of subcases of depression and 82 other subjects. The age distribution of each diagnostic group was similar. More women were present among the cases compared to subcases of depression and non-cases. Most subjects were in social class 3, 4 and 5. Around 40% of both cases and subcases of depression were living alone and this was higher than for non-cases. Stressful events, such as bereavement, illness in the family, house break-ins, family disputes in the preceding month, were more often recorded for subjects of the depressive neurosis group.

The overall dropout rate at 5 years was 28% for cases of depression, 30% for subcases of depression and 26% for other non-cases.

Over 5 years, 41 of 120 cases (34%) died, also 12 of 47 subcases of depression (26%) and 9 of 54 non-cases (17%). The risk of dying within 5 years was 2.1 times higher for the cases of depression compared to non-cases (95% CI, 1.1–3.9). Of the surviving and available cases of depression, 54% (43/79) at year 3, 43% (23/53) at year 4 and 54% (25/46) at year 5 had sufficient psychopathology to reach AGE-CAT case levels for some kind of mental illness. A further 18% (14/79) at year 3, 17% (9/53) at year 4 and 9% (4/46) at year 5 had psychopathology at the subcase level of depression.

LONGITUDINAL OUTCOME

The outcome of the cases of depression is given in Figure 1. Forty-six of the cases of depression had a complete 5 year evaluation. Eleven (24%) had sufficient symptoms to reach AGE-CAT case levels at all follow-up assessments. Another 11 (24%) reached AGE-CAT case levels at two of the three follow-up waves, whilst the other 14 (33%) reached case levels at only one. The remaining 10 (22%) never reached AGE-CAT case levels at any of the 3, 4 and 5 year follow-up waves. The AGE-CAT diagnosis at any follow-up wave was depression in 70% of the interviews. Those (30%) who were diagnosed as another type of case, other than depression, had a high depression score. Only one of the year 3 cases of depression was on antidepressant treatment and none of year 4 and 5 cases of depression were on antidepressants at the time of their interviews.

Outcome of the Subcases of Depression

As many as 16 (34%) of the 47 subcases of depression were AGE-CAT cases at year 3, most of them cases of depression. Their 5 year outcome is given in Figure 2.

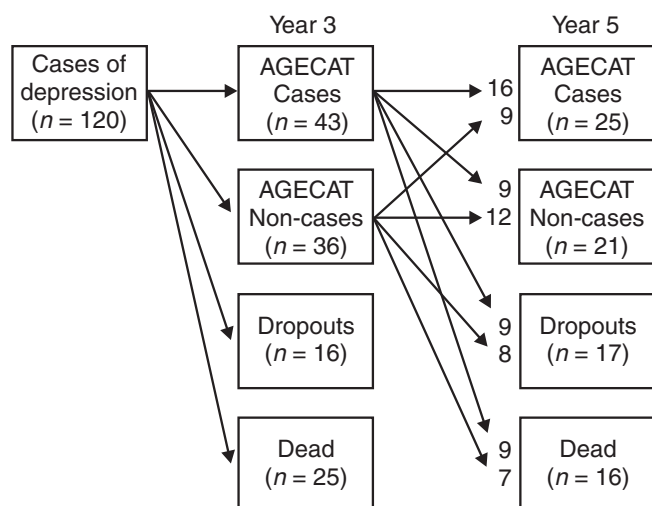


Figure 1 Outcome of the cases of depression (from ref. 34, with permission)

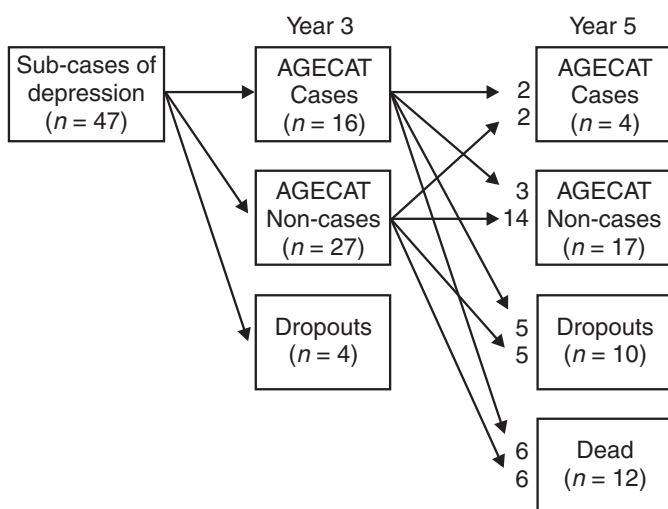


Figure 2 Outcome of the subcases of depression (from ref. 34, with permission).

Organicity and Outcome of the Cases of Depression

In all, 30 (25%) of the 120 cases of depression had co-morbid organic symptoms (AGECAT levels 1–2) at the index assessment. By year 5, nine (30%) had died and seven (23%) had dropped out. Of the remaining 14, 12 (40%) were AGECAT cases and two were non-cases. Seven (6%) of the 120 cases of depression (15% of the surviving cases) had become organic cases at the 5 year follow-up. Five of them had already had some organic symptoms (AGECAT levels 1 or 2) at the start of the study.

Predictors of Outcome

Univariate analyses were done to see which index variables (age, gender, marital status, social class, stressful events, living

conditions, social support, physical illness, depression type (DP and DN), depression score and psychiatric co-morbidity) were predictive of a poor outcome for the depression cases that had survived. Anxiety caseness and a depression score of 30 were particularly predictive of a poor outcome in this sample. Those aged under 75 years did better than those 75 years and older.

A prospective follow-up of the cases of depressive disorders at shorter intervals is the strength of this study. Its findings highlight the importance of the mental health needs of this population. Further studies are needed to discover whether drug treatments or other psychosocial interventions can alleviate depression in elderly people living in the community.

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Longitudinal Studies of Mood Disorders in the USA

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Depressive disorders in late life tend to recur or persist and change the course of a person's life through time. For this reason, the assessment of depression through time necessitates an accurate understanding of the longitudinal course of depressive disorders. For this reason, when the Psychobiology of Depression Study Group began their extensive study of inpatient and outpatient persons in adulthood (but not including old age), they incorporated an extensive longitudinal study into their methodology. Results from this study revealed that 1 year following identification of an index episode of major depression, 50% of the cohort 59 years of age and younger had recovered, but the annual rate of recovery decreased to 28% by the second year and to 22% by the third year. These investigators determined that recovery from an index episode of major depression was most likely to occur within the year following identification of the episode. Among those individuals who do recover, 24% suffered a relapse within 3 months of the recovery; 16% of the individuals who identified suffering from an index episode remained ill throughout the first year. The cohort has now been followed for 15 years (original age of 18–59 with the current cohort now between 33–74); 85% of those who recovered from the index episode relapsed at least once over the 15 years and 58% of persons who recovered and remained well for 5 years relapsed over the next 10 years. Female gender, a longer index episode of depression, more prior episodes and never marrying all contributed to an increased likelihood of relapse.

There has been no equivalent study of the outcome of late-life depression within the USA that has currently been reported in the

literature. However, an ongoing study by Duke University investigators has found that rates of chronicity, recovery and relapse are virtually identical to those reported by Keller and colleagues for younger age groups. The remainder of studies in North America have concentrated primarily upon outcome in controlled treatment trials. The Pittsburgh group found that over 3 years, subjects treated with optimal doses of antidepressant medications relapsed at a rate of between 30% and 40% over 3 years.

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Outcome of Depression in Finland

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In the early 1980s, an epidemiological study of depression was carried out among the population aged 60 years or over ($n = 1529$) living in the municipality of Ähtäri in south-western Finland. DSM-III criteria were used to diagnose depression in interviews and examinations and 264 depressed persons (91 men and 173 women) were discovered; 42 met the criteria for major depression, 199 for dysthymic disorder, 21 for atypical depression and two for cyclothymic disorder.

The depressed persons were intensively treated in a primary care setting for about 2 years after the epidemiological study. The treatment consisted of individual psychological support by a general practitioner, antidepressive medication, counselling about nutrition and physical exercise and social support from the families, relatives, neighbours and home care personnel. Later, a less intensive treatment schedule, consisting of psychological support by a general practitioner and antidepressive medication, was arranged for those who had not recovered.

1 YEAR AND 5 YEAR CLINICAL OUTCOMES

Nearly half of both major depressive and dysthymic patients were non-depressed after 1 year (Table 1). The proportion of depressed subjects tended to be higher among the dysthymic patients, while the proportions of demented subjects and deaths tended to be higher among the major depressive patients. After 5 years, the proportion of recoveries was higher among the dysthymic patients, and the death rate was higher among the major depressive patients. Every fourth subject was depressed in both groups.

FACTORS RELATED TO RELAPSES OR LONG-TERM COURSE

Major depression had a definite tendency for a relapsing course during the 5 year follow-up, even without any special stressors in life or physical illnesses after recovery. The depressed patients who developed a physical disease and whose physical health deteriorated during the treatment had a high risk for non-recovery and a long-term course of depression. Many of these patients had suffered from poor self-appreciation and diurnal variation of symptoms at the onset of the treatment.

The results support the following proposals for clinical practice. Major depressive patients should be followed up after recovery in order to detect their possible relapse and to increase the probability of recovery. Intensive antidepressant and psychotherapeutic treatment and adequate treatment of physical diseases should be arranged for depressed patients who develop a physical disease or whose somatic condition deteriorates due to a previous physical disease. Cooperation between psychiatrists and general practitioners is needed in the above two cases.

MORTALITY

Major depressive patients had a high death rate, which was not explained by their poor physical health. These results suggest that there may be biological factors associated with major depression that increase the risk of death or the risk of the development of physical diseases leading to death. The mortality of dysthymic patients was

Table 1 One year and 5 year outcomes of major depressive and dysthymic older patients treated in primary health care

	1 Year outcome		5 Year outcome	
	<i>n</i>	(%)	<i>n</i>	(%)
Major depressive patients				
Non-depressed	19	(45)	5	(12)
Depressed	11	(26)	11	(26)
Demented	6	(14)	5	(12)
Dead	6	(14)	19	(45)
Refused to participate			2	(5)
Total	42	(100)	42	(100)
Dysthymic patients				
Non-depressed	79	(40)	71	(36)
Depressed	91	(45)	52	(26)
Demented	6	(3)	18	(9)
Dead	20	(10)	50	(25)
Refused to participate	3	(2)	8	(4)
Total	199	(100)	199	(100)

also higher than that of non-depressed persons, but their high mortality was explained by the high number of physical diseases.

Longstanding depression was a predictor for high mortality, independently of the physical diseases present at the onset of the treatment of depression. The physical diseases that occur during the treatment and predict a longstanding course may explain the high death rate seen here.

These results also underline the need for intensive and adequate treatment of depression and physical diseases in depressed older persons.

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Physical Illness and Depression

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EPIDEMIOLOGY

The prevalence of depression in the general population worldwide is usually found to be 3–8%^{1–3}. The prevalence of major depression has been shown to be no higher in the elderly than the young, although these findings do not allow for the comorbidity of physical illnesses or dementias⁴. Subthreshold or minor depressions have many different names and definitions, thus causing widely differing prevalence rates to be quoted. Categorical definitions of depression do not fit well with the range of symptoms and severity seen in normal clinical practice. However, it is generally accepted that the burden of depression among the elderly is high and an accepted measure of diagnosis is necessary to allow communication with patients, relatives and professional colleagues.

“Caseness” can be considered to be the severity of depression at which the majority of professionals would consider some form of intervention appropriate. Prevalence of this degree of depression is reported as 10–15% of the elderly in the community^{5–6}, 15–30% of those attending primary care facilities^{7–8}, 15–50% of those in hospital^{8–10} and 30–40% of those in institutional care^{8,11}.

DIAGNOSIS

Depression cannot be diagnosed unless it is first considered a possibility, neither will it be appropriately treated unless it is considered pathological. Depression may be missed when too much emphasis is placed on the presenting complaints of, for example, lethargy, anorexia or pain¹². Depression and feelings of worthlessness may cause failure to complain of symptoms of physical illness or to ask for help. It may cause non-compliance with medication and other treatments, self-neglect or non-attendance at clinics. Alternatively, the lowering of self-esteem and decreased ability to cope can lead to increased attendance at clinics.

The lack of a concise definition for depression in the elderly makes the establishment of validity a difficult task, which can only be examined by longitudinal follow-up of patients to see what happens to their symptoms¹³. Somatic symptoms, e.g. lack of energy, poor concentration and weight loss, may be due to the physical illness or ageing, not depression; even experienced clinicians may have difficulty attributing such symptoms to physical or psychiatric causes. Even feelings of life not being worth living and wishing to die are not always associated with depressed mood; poor subjective health, disability, pain, sensory impairment and living in an institution have been shown to be associated factors in the absence of depressive illness¹⁴.

The elderly tend not to admit to feelings of depression and relatives may be unaware of the condition¹⁵. Somatization, “the tendency to experience and communicate somatic distress and somatic symptoms unaccounted for by relevant pathological findings, to attribute them to physical illness and to seek medical help for them”¹⁶ is increasingly recognized. Somatization can still occur in those with genuine physical illness. The somatic symptoms of depression are similar to those of a chronic illness, such as cancer, and it must be remembered that depression and physical illness often coexist¹⁷.

Hypochondriasis is a recognized symptom of depression in the elderly population¹⁸. However, in this age group, rigorous steps must be undertaken to exclude physical problems before ascribing symptoms to hypochondriasis or somatization^{17,19}. That such patients are depressed is inferred from their good response to standard treatments for depression²⁰.

The 1991 NIH Consensus Statement on diagnosis and treatment of depression in late life concluded: “What makes depression in the elderly so insidious is that neither the victim nor the health provider may recognize its symptoms in the context of the multiple physical problems of many elderly people”. DSM-4 allows somatic symptoms to be counted towards the diagnosis of depression if there is any possibility of psychological aetiology, a more inclusive and accurate means of diagnosis than previously.

MORBIDITY AND MORTALITY

Psychiatric morbidity in hospitals is higher than in the general population. Surveys of wards and clinics do not completely establish an association between psychiatric and physical morbidity because they may be biased for selective referral patterns: psychological symptoms can lead to help-seeking behaviour for physical illness in an individual who had previously been able to tolerate his/her physical problems. Similarly, they may influence a GP on whether or not to refer to hospital. Stress may be as important in triggering help-seeking behaviour as in triggering actual illness. In addition to the degree of distress, many other factors determine whether or not an individual will seek help, including religious and social values, socioeconomic background and personality.

Affective disorder in the elderly is strongly associated with physical ill-health¹⁹: “whether or not such an illness has a direct aetiological relationship to the affective disorder, its practical importance must be considered, for it is bound to influence the course and outcome of the psychiatric condition”. Other studies have found that depression leads to increased mortality^{21–23} over and above age effects, the prognosis worsening with severity of

depression. Depression in the elderly may be due partly to a biological ageing process, which would directly increase mortality and morbidity^{20,24}. Burvill and Hall²⁵ showed increased mortality in depressed elderly patients ($n=103$, age 60+) followed for 5 years if they were aged 75+, had impaired mobility or showed poor recovery with residual symptoms or chronicity. There were two peaks of increased mortality, one early in the disease and one late. Cardiovascular or pulmonary disease and malignancies were the predominant causes of death. The results are similar to those of Murphy *et al.*'s 4 year follow-up²⁶, who also postulate that increased mortality seen in the depressed elderly (especially the men) was not due to differences in physical health alone. They suggest:

1. Inadequate treatment of the depression, leading to cardiovascular complications from the antidepressant but no benefit to the patient; the depression itself can also provoke cardiac death, especially in men.
2. "Subintentional suicide" in those who "turn their faces to the wall".
3. Residual depressive invalidism, causing poor nutrition and decreased mobility; with attendant complications of susceptibility to infection, fractures, bedsores, etc., all contribute to the increased mortality.

Mortality in hospital was found to be significantly higher in those depressed but over 30% of those discharged had died within, on average, 5 months, whether depressed or not²⁷. The authors also noted that survivors with depression consumed more healthcare resources than did the non-depressed survivors. Among the depressed elderly, 40% have chronic poor physical health²⁸; they use and need more medical services^{29,30} than the non-depressed, but also use fewer social and recreational services²⁹. An association has been found between poor mental health and subsequent physical disease, suggesting that positive mental health may significantly retard the decline in physical health with increasing age³¹.

Physical illness affects the capacity for independent living, resulting in altered relationships with others, lowered self-esteem and vulnerability to depression. Serious illness may be seen by some as an unpleasant reminder of mortality, bringing apprehension and fear. Continuing physical illness is a poor prognostic factor for depression, although whether this is due to a biological relationship or to the psychological strain of being ill is uncertain.

Mortality in acute medical inpatients is significantly higher in those with associated depression, although the direction of causality is not established, e.g. Silverstone³² followed consecutive admissions for myocardial infarction, subarachnoid haemorrhage, pulmonary embolus or upper gastrointestinal haemorrhage for 28 days post-admission; 34% were depressed and 47% of these had life-threatening complications or died, compared with 10% of those not depressed.

PHYSICAL ILLNESS AND DEPRESSION

Mood disturbance can result from structural brain disease, alterations in neurotransmitter concentration or activity caused by drugs or biochemical disturbance. These affective symptoms may present during a physical illness or be the initial symptom of an otherwise occult physical disorder. Depression may be:

1. The result of an illness, especially a painful or disabling one.
2. Iatrogenic, e.g. the result of steroid treatment.
3. A symptom of the physical illness, e.g. hypothyroidism.
4. An aetiological factor, e.g. alcohol abuse secondary to depression.

5. A depressed patient adopting the sick role as a coping mechanism.
6. A common aetiological factor, such as bereavement, may cause both depression and physical illness.
7. Coincidental.

The elderly are particularly susceptible to the side effects of drug treatment³³, especially as they are often subject to polypharmacy^{34,35}. Patients with drug-induced depression often have a past or family history of depression and the drug may have precipitated the disease by affecting the levels of available neurotransmitters³⁴. Depression can often be alleviated by cessation of the drug, although some patients will also require antidepressant treatment. The combination of a susceptible patient and a depressogenic drug may precipitate a depression sufficiently severe to lead to suicide³⁴.

Subjective rating of general health has been shown to be independently associated with depression in the elderly, including the very old³⁶⁻³⁸. This can lead to presentation at primary care or emergency facilities, unnecessary investigations and risk of iatrogenic disease, and lower quality of life. Recognition and treatment of the depression may lead to improvement in the patient's subjective perception of his/her health.

Physically ill depressed patients are more likely to be admitted to hospital than those who are not depressed³⁹. Depressed patients have higher use of all categories of medical care, including admissions, laboratory tests and emergency department visits³⁰.

The presence of significant psychiatric disorder has been shown to adversely affect the course of medical admission^{40,41}, affective disorders in particular prolonging length of stay⁴²⁻⁴⁴ (although the study by Ramsay⁴⁵ did not confirm this) and increasing the likelihood of admission to residential care⁴⁶. Psychiatric intervention has been shown to increase recovery rate, reduce duration of stay, reduce the need for residential care after discharge and therefore reduce costs^{47,48}.

ADJUSTMENT DISORDER

Lipowski⁴⁹ has proposed that the subjective significance of an illness and its treatment, e.g. amputation, cancer, combined with the patient's personality and social circumstances, is the key to the psychological response. The variety of physical illnesses found with depression would support this view. Depression may be a reaction to physical problems; it occurs more frequently in those with increasing numbers of medical diagnoses and may be precipitated by developing new physical illnesses^{21,50}. All illnesses except the very trivial involve an element of psychological adjustment. Serious medical illness is likely to be a potent psychological stressor, affecting body image, self-esteem, the sense of identity and the capacity to work and to maintain social, family and marital relationships⁵¹. However, the majority of people adapt their lives to the demands of their illness, maximizing their prospects of recovery and return to previous levels of activity.

In the elderly, physical illness is frequently chronic and may worsen with time. This, combined with the losses suffered by many elderly people, such as loss of status and income on retirement, loss of friends and family by death and the fear of loss of independence and dignity due to the illness itself, can lead to the adjustment disorder merging imperceptibly into a depressive illness. This can be considered secondary depression⁵², the depression following or paralleling a life-threatening or incapacitating medical illness. However, the prevalence of this type of depression is unknown, as it is difficult to differentiate from depression related to other stressors and previous history. Patients with this type of depression tend to have fewer suicidal thoughts but have more feelings of helplessness, pessimism and anxiety⁵³.

SOMATIZED DEPRESSION

This is more common than medical illness presenting as depression, especially in the current generation of elderly, who tend to somatize their psychological symptoms, having been brought up in a society which did not encourage the expression of emotion. Somatic symptoms in the elderly may represent physical illness, depression or emotional responses to physical illness. The somatic symptoms will need investigation, but depression, if suspected, should be treated.

Pseudodementia is a specific type of masked depression. One of the most important differentiating factors is that the severity of cognitive impairment fluctuates in depressed patients, remaining constant or worsening in the evenings in dementia. Depressed patients tend not to try to succeed in tasks, giving up with "I don't know" or "I can't". Demented patients will try, delighting in success but possibly becoming very distressed by failure—the so-called catastrophic reaction. Biological symptoms of appetite and weight loss, sleep disturbance and headache are typical of depression and not dementia. However, depression and dementia can coexist and the differentiation of the two conditions is not always easy. If in doubt, a trial of antidepressant treatment will help elucidate the diagnosis. The pathognomic symptoms of masked or somatized depression include:

1. Diurnal variation (symptoms usually worse in the mornings).
2. Mild impairment of cognitive processes and concentration.
3. Dysthymic mood changes.
4. Fatigue, feeling tired, lack of energy.
5. Sleep disturbance (waking up early and being unable to get back to sleep).
6. An anxious sense of failure or of "impending disaster".

LIAISON

Consultation-liaison psychiatry is becoming well established as an important specialty within general hospitals on both sides of the Atlantic. In an ideal situation, psychiatrists would attend ward rounds in the general hospital, particularly on rehabilitation wards, where prevalence of depression is high and the effect on delayed discharge well documented⁴²⁻⁴⁴. However, restricted resources prevent this: psychiatric morbidity is too high for a psychiatrist to see all the patients affected—his main role should be educational⁵⁴, only taking an active part in the management of more difficult cases. In many areas there is increasing development of specialist liaison nurses who are able to advise on diagnosis and treatment, reducing delay before assessments. The liaison nurse is likely to be more permanent than junior doctors on rotation and can often help a patient who refuses to see a psychiatrist or whose physician refuses psychiatric referral⁵⁵. Liaison nurses can educate general nurses in the recognition of psychiatric disorder and can in turn encourage junior medical staff to institute appropriate referral or treatment. Their development and use has been compared with that of community psychiatric nurses⁵⁶.

The most common reasons for requesting a psychiatric consultation in a general hospital are⁵⁷:

1. Diagnostic uncertainty.
2. Recognition of a gross psychiatric disorder.
3. Excessive emotional reactions, e.g. fear, anger, depression.
4. A patient's deviant behaviour disturbing ward or medical procedures.
5. Delayed convalescence, i.e. disability incompatible with observed pathology, relapse on mention of discharge.
6. Crisis in the doctor-patient relationship (e.g. refusing consent!).

7. Patient's admission of serious psychosocial difficulties.
8. Selection and/or preparation of patients, e.g. pre-transplant, cosmetic surgery.

It can be seen from the above list that the depressed patient, sitting quiet and withdrawn on the ward or in a home, may not be referred for a psychiatric opinion. In practice, only about 2% of geriatric patients are referred to the liaison services⁵⁸⁻⁶⁰. Suggested reasons for the discrepancy between liaison rate and psychiatric morbidity are^{58,61}:

1. High prevalence of transient self-limiting psychiatric disease.
2. Physician's failure to recognize psychiatric disease⁶². Many studies have highlighted the unrecognized psychiatric problems on medical wards⁶¹ and the underdiagnosis of major depression in particular has been well documented⁶³.
3. Medical and nursing staff may actively avoid questioning for psychological problems, due to fear of precipitating emotional distress with which they have not been trained to deal.
4. The low priority of psychiatric disease compared to physical, especially in a busy medical ward with acutely ill patients.
5. Poor access to, or dissatisfaction with, psychiatric services.
6. Physician resistance to psychiatric consultation⁴⁷, due to stigma or underestimating the severity or the potential for treatment.

Use of screening scales is appropriate for assessing depression in the physically ill: it is common, can be a difficult diagnosis and has significant morbidity and mortality if untreated. Care must be taken to differentiate between short-lived adjustment disorders, occurring as a reaction to the admission itself, or the crisis which precipitated it. Diagnosis in acute admissions should therefore include enquiry into symptoms before admission. If none can be elicited, the patient should be reassessed at a later date, either during rehabilitation or after discharge.

Screening scales serve a dual function⁶⁴⁻⁶⁶—they identify patients in need of further assessment and also serve as an educational tool if given by general staff during routine admission procedures. They emphasize the associated symptoms and signs of depression in the elderly, who may not show depressed affect and will deny feeling sad. It is important, however, that education about the treatment of depression goes hand in hand with education to recognize it, or the liaison services will be swamped.

TREATMENT

The fact that one can intuitively "understand" why the physically-ill elderly are depressed does not mean that it should be accepted as normal and treatment not attempted. Continuing physical illness is recognized as both a precipitant of depression and a poor prognostic factor, yet despite this, ~80% of elderly general hospital patients are not depressed^{42,67,68}. Successful coping mechanisms can prevent the emergence of clinical depression. Even in the terminal patient, depression or dysphoria can be relieved by euphorants, such as oral or parenteral opiates, reducing the distress of patient and relatives⁶⁹ without significantly reducing the length of life remaining to the patient.

It is important that the diagnosis is not missed, as this condition usually responds well to treatment, at least initially, thus improving quality of life. Follow-up and early treatment of any relapses will further improve the prognosis. Increased self-esteem and ability to cope will reduce demand on families and possibly on services.

Even when the correct diagnosis is made, the depression may not be treated adequately, if at all: physically ill patients who are also depressed are more likely to be assigned to the "not to be resuscitated" group, compared with those elderly who are not

depressed. The lack of effort and motivation caused by depression may be regarded as “not trying” or “giving up” by nurses and rehabilitation staff, who withdraw from this group of patients for more emotionally rewarding non-depressed elderly subjects on the same ward.

The elderly as a group have approximately twice as many adverse drug reactions as younger adults⁷⁰. They are frequently already on polypharmacy, so drug interactions are a real possibility. It is therefore important that physicians are advised and supported by the psychiatric services in the use of safe, well-tolerated antidepressants and other treatments to reduce the impact of this disease on both the individual and society.

Electroconvulsive therapy (ECT) is an effective treatment of depression^{71,72}. The response to treatment in older people is better than in the young^{73,74}. With the increasing safety of anaesthesia, very few patients, even those with severe physical disease, are unable to tolerate a course of treatment. It is more rapid in effect than medication alone, but the improvement is rarely sustained unless antidepressants are also given to prevent relapse.

A review of psychological treatments in chronic illness⁷⁵ found very little empirical evidence of benefit when therapeutic interventions were applied indiscriminately. However, some evidence was found to show benefit in patients with somatization disorders rather than physical illness, and also that brief interventions following the onset of acute physical illness reduced longer-term psychological morbidity. A suggested general approach to the treatment of physically ill depressed patients is:

1. Investigate and give appropriate treatment to all physical problems, either curative treatment or to minimize persistent morbidity. Explain the illness, treatment and prognosis to the patient in as much detail as he/she wishes. Make sure the explanation is understood and repeat as often as necessary.
2. Give general social support, e.g. home help services, financial assistance if relevant, or residential or nursing home care.
3. Give psychological support—encouragement, continued interest in the patient, e.g. outpatient follow-up. Support groups are often beneficial for chronic conditions such as rheumatoid arthritis, Parkinson's disease, etc.
4. Consider antidepressant therapy if the symptoms are sufficiently severe that they would be considered to warrant medication if seen in a patient without physical problems. Monitor the response to treatment; this may take 7–8 weeks⁷⁶. If no response is seen to a therapeutic dosage of antidepressant, consider a trial of an alternative antidepressant or adjunctive treatment, or specialist referral.

PROGNOSIS

Little is known of the prognosis of psychiatric disorder identified in the medical setting⁷⁷, except that concomitant physical illness is a poor prognostic factor. Psychiatric disturbance often persists^{78–80}, especially in patients with a previous history of psychiatric disorder. Those with affective disorder on admission have increased mortality and make greater demands on medical, social and psychiatric services^{77,81}.

In one series of consecutive acute medical admissions⁸⁰, fewer than 45% of those patients with concomitant depression had received antidepressants at all, 20% had been given benzodiazepines, and less than 25% had been treated for more than one week. The authors concluded that an effective treatment for depression in elderly patients needed to be found, with widespread education of geriatricians in the diagnosis and treatment of depression.

In psychiatric patients, relapse has been linked with super-vening physical illness⁸². The presence or development of cerebral

or any other irreversible physical disorder indicates poor future mental health in the great majority of patients, as well as the likelihood of early death⁸³.

The prognosis of the physical illness, for both morbidity and mortality, is also inextricably linked with that of the depression. Increased mortality from physical illness, especially cardiovascular disease, has been reported^{84,85}. This excess of deaths is significantly associated with groups who have been only partially treated, e.g. have not responded to antidepressants⁸⁶, especially in older men⁸⁷.

Explanations for the apparent association of physical illness with poor treatment outcome might include⁸⁸:

1. Age as a confounding factor—most of the trials of treatment in the physically ill are in the elderly.
2. Medically ill patients may be given inadequate doses of antidepressants because of problems with side effects or over-cautious physicians.
3. Different subtypes of depression exist, some medication-responsive, some not. Organic mood disorder (depression induced by physical illness or a specific organic factor) has been shown to have a worse prognosis at 4 year follow-up⁸⁹.

The prognosis of depression in the physically ill elderly is therefore dependent on accurate diagnosis, intensive treatment, follow-up and early treatment of any relapses^{82,84,90}. Increased self-esteem and ability to cope will reduce demand on families and possibly on services. No studies have yet convincingly identified predictors of response in physically ill populations, although pre-existing depression prior to admission with physical problems appears to predict persistent depression, rather than if the depression develops in hospital^{78,91}.

SUICIDE

Suicide is the most dramatic of poor outcomes, and the elderly are over-represented in suicide statistics. Although the elderly are less likely to attempt suicide, they are more likely to complete it^{92,93}. The presence of physical illness, especially if associated with chronic pain and disability, increases the risk; elderly men living alone are at the highest risk. The individual's adjustment to ill-health and his associated feelings of hopelessness and demoralization are obviously important⁹⁴. Many elderly suicide victims are suffering from their first episode of major depression, which is typically only moderately severe but the diagnosis is missed⁹⁵, and the potential for recovery following intervention therefore lost.

Depression has been linked to decreased compliance and to voluntary refusal of life-saving essential medical treatments^{96,97}. This may reflect either conscious or unconscious suicidal motivation.

SERVICE IMPLICATIONS

It is important to note that the aging population itself is growing older, with large numbers of very old individuals. It is this old-old group who have the highest physical and psychiatric morbidity and who make the greatest demands on services.

Undergraduate teaching programmes must be tightly integrated in order for students to develop a holistic approach to the elderly, together with an understanding of the psychosocial and economic factors that will affect presentation and treatment. Joint post-graduate meetings between the two specialties are becoming more common and should be encouraged, each maintaining their separate identities and training but working closely together in clinical practice.

Interdisciplinary research continues to grow in amount, but is mostly directed at psychiatric problems on medical wards. Little seems to be researched in the other direction, medical illnesses on psychiatric wards. All research is obviously relevant to both specialties and in practice helps to strengthen links between them.

CONCLUSION

Depression in the elderly physically ill can present a difficult problem of diagnosis. To avoid the increased morbidity and mortality of untreated depression, it is necessary to be aware of the possibility of depression causing somatic symptoms, the physical illness causing depressive symptoms, or both conditions coexisting, and then to treat the depression effectively. Close liaison between psychogeriatric and geriatric teams is thus very important if this vulnerable group is to receive the correct diagnosis and treatment.

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Physical Illness and Depression: a Number of Conundrums

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One of the clinical conundrums in modern medicine is that psychiatric patients complain frequently in somatic terms and yet *truly* suffer from an excess of physical illness¹. In a review conducted as part of a European Science Foundation Study (EMRC), Hafner and Bickel² concluded that "Studies of mortality in mental patients have shown that... these patients still have an excess risk for natural causes of death which is not restricted to patients or to certain diagnostic groups... However, the evidence for *specific* associations between psychiatric diagnosis and natural causes of death is not yet conclusive". In a commentary, Rorsman³ said that two out of three major *longitudinal* studies indicate, at least in men, that mental illness strongly affects the risk of dying from natural causes. Murphy *et al.*⁴, from the Stirling County study, found death to be significantly associated with affective, not physical disorder, and depression, not anxiety. Rorsman *et al.*⁵, from the Lundby study, found that psychiatric patients have a significantly increased natural death risk, and *untreated* psychiatric males in particular.

An issue of the *International Journal of Geriatric Psychiatry*⁶ dealt with physical illness and depression in the elderly. Burvill, from Australia, pointed out that physical illness worsens the prognosis of depressive illness in the elderly. Lindsay, from the Guy's/Age Concern Survey in the UK, found that 70% of depressed subjects reported one or more serious physical problems. Sadavoy *et al.*, from Canada, found that about 75% of the elderly with chronic physical illness had cognitive impairment and 35% were depressed. There was a significant correlation between cognitive deficit and depression.

Eastwood and Corbin⁷, in a review of the connection between depression and physical illness in the elderly, addressed another conundrum. While physical disease increases with age, depression may not do so. In community surveys of the elderly, fewer than 25% are disease-free and over 50% have at least one activity-limiting disorder⁸. While the findings are disputed, depressive illness apparently declines with age, while depressive symptoms increase. Snowden⁹ argued that depressive symptoms and syndromes are difficult to distinguish in the medically ill. He thought that, since conditions which significantly correlate with depression, such as dementia, physical disability, physical illness, bereavement and so on, increase with age, then so must depression. Recently, Mann¹⁰ argued that, "if other depressive, diagnostic terms are included—'minor depression', 'subthreshold syndrome' or 'depressive symptoms'—then the total rate of depression is, in fact, higher than in the younger age groups". The truth probably lies in some complex multivariate relationship. Notwithstanding, there are some fascinating and relatively direct relationships, such as stroke causing depression¹¹ and grief causing increased coronary heart disease¹². Fascinatingly, Glassman and Shapiro¹³ consider that we have reached the point where we can state that depression is an independent risk factor for

coronary heart disease. While taking this as an interesting postulate, it has to be remembered that atherosclerosis could be a cause of both depression and heart disease. At this stage we do not know whether intervention with antidepressants would reduce the risk of depression on heart disease.

Finally, as Hafner and Bickel suggest, prospective studies with disease registers will help sort out general and specific relationships and direct and indirect risk factors, and help confirm Rorsman's statement, that this all means that psychiatry is a branch of medicine.

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Depression after Stroke

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It is now accepted that stroke patients have high rates of all types of depressive disorder. However, estimates of the prevalence of depression within the first month of stroke vary greatly^{1,2}, according to the type of measure used and the way the sample was derived. The consensus seems to be that 20–25% of patients after stroke will suffer a major depressive disorder within the first month³. Rates of depression later on after stroke are less certain, because of patient attrition in studies. Depression remits in some patients, while in others it persists: one study reported that 50% of those depressed within 3 weeks of stroke remained depressed 1 year later⁴. Depression not only affects quality of life: patients with depression after stroke may be at greater risk of mortality⁵, cognitive impairment⁶ and poorer functional or social recovery^{7,8}.

There are claims that post-stroke depression is a distinct subtype⁹. There is little unequivocal evidence to support this claim, since psychological and biological symptoms reported by patients are found in depression seen in non-stroke patients¹⁰. What does appear to be distinctive is the increased prevalence of persistent crying (emotionality) among stroke patients. A small number of patients suffer *pathological laughing and crying*, a syndrome which is probably neurological in origin, in which emotional expression arises after minor provocation that often appears meaningless¹¹. A more common syndrome, *emotionalism*, is also characterized by increased tearfulness, but is more complex. The emotional episodes are provoked by meaningful stimuli, but the crying is characterized by a lack of warning and control. There appears to be a psychological component to its origin¹². *Emotionalism* is associated with an increased risk of depression¹³, but patients with emotionalism may be at greater risk of psychological problems not explained by concurrent depression¹⁴. This syndrome is probably under-recognized.

Reaching a diagnosis of depression after stroke can be complicated by the presence of problems such as communicative or cognitive impairment¹⁵. Patients with expressive communication problems can often be assessed by the careful use of closed questions. The assessment of those with receptive communication problems or significant cognitive impairment is much more complex: a non-language-based assessment of depression shows promise but is insufficiently reliable in its present form for accurate diagnosis¹⁶. The diagnosis of depression might also be confused by facial palsy and a disturbance of speech prosody (rhythm in speech), both of which are relatively common after stroke and which give the patient the appearance of a person with depression¹⁷. The dexamethasone suppression test is not sufficiently sensitive to be used as a diagnostic tool¹⁸.

The high rate of depression reported in some stroke research has led to the suggestion that the neurological damage is a key factor in its aetiology¹⁹. As a result, many studies have attempted to link depression after stroke with lesion location. A series of

studies proposed, first, that patients with left hemisphere lesions were at greater risk of depression²⁰, and later, that those with left anterior lesions were at most risk^{21,22}. Other researchers^{23–25} have not replicated these findings, suggesting that patient sampling and the timing of assessment might explain the differences.

Even if lesion location is associated with greater risk of depression, the context of this relationship is important. First, it is clear that stroke patients with all sorts of lesions can suffer depression⁹, so factors other than lesion location must also be at work. Second, stroke location is extremely varied²⁶, so those with any particular lesion (such as left anterior lesions) will be a minority of patients, making the attributable risk due to any one type of lesion small. Last, although the rate of depression in stroke patients is higher than in age-matched non-stroke controls, it is about the same rate as in patients with non-neurological disabling illness^{17,27}, suggesting that non-neurological factors are as important. Relevant non-neurological aetiological factors are likely to include the threat of disability and a sense of loss.

That psychosocial factors are likely to be relevant to both the onset and persistence of depression has been illustrated in several studies. For example, one study found that depression at 4 months after stroke onset was commoner among those with greater disability, those who were divorced and those with higher pre-stroke alcohol intake²⁴. Depression is more likely in those patients who perceive their stroke as a greater threat, and in those who have fewer psychological resources to deal with that threat.

Patients with depression after stroke might be considered for pharmacological or psychological treatments. Two small trials showed beneficial effects of antidepressants, although both studies had high rates of patient dropout^{28,29}. The evidence for treating emotionalism with antidepressants is rather stronger—both tricyclics and SSRIs have been shown to reduce the frequency of crying episodes^{30–32}. There is no good trial evidence to draw upon in assessing whether psychological interventions are effective in treating depression after stroke³³.

Some services aim to intervene in an attempt to prevent the onset of depression after stroke. A recent small trial found that patients prescribed mianserin as a prophylactic had greater improvement in depression scores, but the drug did not reduce rates of major depression 6 months after stroke³⁴. A variety of preventive psychosocial interventions have been evaluated in clinical trials. The interventions, including education, leisure therapy and specialist stroke nurse visits, have shown no effect in reducing the prevalence of depression. However, many of the trials are small and imperfectly designed, so the conclusion should be lack of evidence, rather than evidence of no effect³⁵.

Our own recently completed study suggests that a brief psychological treatment (problem-solving therapy) may be beneficial^{36,37}. Patients who received therapy visits from a community

psychiatric nurse had lower rates of depression 6 and 12 months after stroke than those in the treatment-as-usual group, and lower scores on a measure of psychological distress at 12 months. This finding is encouraging, since it shows that a brief, structured psychological intervention is beneficial to patients after stroke, albeit in a sample selected to participate in a clinical trial. There are disadvantages to psychological management: it may be difficult to implement in patients with significant speech and cognitive impairment³⁸, and some patients find psychological treatments unacceptable, both before and after the treatment has started.

In summary, depression after stroke is common, and its causes are probably multiple—biological, psychological and social—as is the case in other physical illnesses. The evidence for benefit from antidepressant drugs is surprisingly poor, considering their problematic side effects and how widely they are prescribed. The potential for psychological therapies has been undervalued, which is a deficit that badly needs correcting. Pending further research, clinicians will need to rely on evidence from other areas of physical medicine to inform their treatments.

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Treatment of Depression in Older People with Physical Disability

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THE CASE FOR TREATMENT

As discussed elsewhere, depression is the most common mental disorder in the over-65s, with a prevalence of 13–16%^{1,2}. It is a serious disorder, associated with profound decrease in quality of life³, suicide⁴, non-suicidal excess mortality unexplained by physical disorder⁵, and excess health and social service use not explained by disability^{6,7}. Depression in the elderly also has a substantial financial impact, costing community health and social services in the UK in excess of £1 billion/year in depression-dependent service use⁸.

Depression in the elderly therefore has a serious impact on the people suffering from it, their families, and health and social services. Despite this, it is a consistent finding that few older people with depression receive appropriate treatment from primary or secondary care services. Only 10–20% of cases of depression are prescribed antidepressants^{7,9,10}, with no evidence of their receiving non-drug treatment instead. The reasons for this lack of appropriate action is unclear, some implicating low GP recognition¹¹ and some a lack of action when depression is found¹². Whatever the mechanism, there are clear discontinuities on the path from contact, through recognition to action¹³.

Older people with physical illness or disablement are a high-risk group for the development of depression¹⁴. One particular high-risk group consists of those maintained at home, receiving social service home care; 26% of these have clinical depression¹⁵ and, adjusting for age and gender, they have twice the prevalence of depression of the general elderly population, with a four-fold excess of the most severe forms⁷. One possible determinant of therapeutic inactivity may be a perception that depression is untreatable in frail older people, and an important element in clinical behaviour change is evidence of the effectiveness of intervention. Meta-analyses suggest that antidepressants have efficacy in the treatment of depression in those with a variety of physical illnesses¹⁶, with the same sort of effect sizes as those observed in the physically well. However, the evidence for the effectiveness of treatment for depression in the disabled elderly is sparse, since they are often systematically excluded from drug trials¹⁷. We therefore completed a randomized controlled trial (RCT) to investigate whether depression in home-care clients was treatable by community old age psychiatric services¹⁸.

THE EFFECTIVENESS OF OLD AGE PSYCHIATRIC COMMUNITY TEAM INTERVENTION

Sixty-nine cases of depression were identified by screening the home-care population and randomly allocating them to treatment as usual by their GP, or to treatment by the local old age psychiatric community team, with blind follow-up at 6 months. There was a powerful treatment effect, with 58% of the intervention group recovering, compared with only 25% of the control group (adjusted odds ratio 9.0 [95% CI, 2.1–41.5]). The intervention was pragmatic, involving the multidisciplinary team formulating an individualized management plan and this being implemented by a research worker working as a generic team member. Analyses were carried out on an intention-to-treat basis.

This study's results suggest that therapeutic nihilism, based on an assumed poor response to treatment in the disabled elderly, may not be justified. There are similarly encouraging data for the general population of older adults with depression from GP practice-based community psychiatric nurse intervention^{19,20} and nurse-based outreach programmes²¹. However, all these interventions are complex and delivered by secondary care services, and are therefore not directly transferable into primary care settings. Given that there may be 500 000 disabled older adults with clinically significant depression in the UK alone at any one time, secondary care intervention for all is not feasible. It would also be unnecessary if depression in the disabled elderly were to be managed successfully by primary healthcare teams. These are questions which require further research.

Elements of Effective Intervention

The dysjunction in the system of care from disorder to recognition to action has been outlined above. What, therefore, does this mean for the formulation of effective interventions for older people with depression, and where might change be focused best to achieve maximum health gain? These questions can be addressed by considering the pathway from depressed state to resolution, using the data we have for disabled elderly home-care recipients.

Figure 1 presents a simple model. In it, the outcome of depression depends on two parameters, the natural history of the disorder and the effectiveness of intervention. The extent to which an intervention is deployed depends on there being both recognition and action. In Figure 1, the data from the home-care studies are applied to a standard population of 100. In the first stage, the current 15% rate of any active management for depression in this population⁷ is applied to divide the group into a "treated" and a "not treated" group. The second stage is to apply the spontaneous recovery rate of 25% to the "not treated" group and the 60% recovery rate from our RCT with active management to the "treated" group¹⁸. When these filters are applied, only

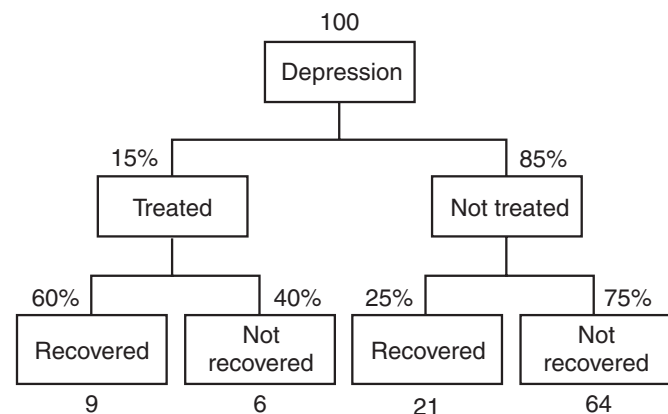


Figure 1

30 of the 100 recover (only nine in the “treated” group and 21 spontaneously recovering in the “not treated” group).

Where, then, should effort be focused to address this situation? The spontaneous recovery rate will be fairly resistant to change and the gain from trying to improve the efficacy of treatment would appear to be relatively limited. For example, an increase in the efficacy from 60% to 80% (an increase unlikely to be possible at present) would only increase the numbers recovering by 3 to 12 in the “treated” group. What is clear from Figure 1 is that the main determinant of the poor population outcome is the low rate of recognition/active management. This would suggest that resources should be focused on increasing the proportion that enter the “treated” group, since there is the greatest scope for improvement at this point and any benefit at this stage will cascade down the system. So, if the proportion “treated” were to be raised by 20% to 35%, the numbers recovering in the “treated” arm would rise to 21 (with 37% recovering overall). If only half of the population of people with depression were identified and treated, then this would rise further to 30 (with 43% recovering overall) and a 75% treatment rate would yield 45 recoveries (51% overall).

CONCLUSIONS

These data demonstrate that, on a population level, there is likely to be far greater health gain from attending to the processes of recognition of depression in the elderly, and of linking this recognition to action, than there is by simply focusing resources on attempting to develop interventions with greater efficacy. This is supported by the emerging evidence base, which endorses the feasibility, acceptability and effectiveness of screening for and treating depression in older adults in the community¹⁸⁻²¹.

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Acute Management of Late-life Depression

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The acute management of late-life depression may require hospitalization, both for accurate diagnosis and for effective treatment. Ambulatory management is frequently favored because of rising hospital costs in a managed care environment. However, the elderly present special challenges that may require that diagnosis and/or treatment be undertaken in a hospital setting. The hospital provides an environment for the monitoring of symptoms for accurate diagnosis and proper personnel for regular and accurate treatment administration. Several factors may interfere with both accurate diagnosis and effective treatment on an ambulatory basis¹, including underlying chronic medical illness, pain, neurodegenerative changes, dementia, adverse life events, inadequate family support, secret self-medication, substance abuse, bereavement, interpersonal conflicts and social isolation of the elderly patient.

Actually, the elderly patient who is cognitively intact may be reliably managed on an ambulatory basis and can be instructed about medication side effects. Similarly, the more impaired elderly patient who has adequate social support for observation and medication management may only need the community support of a visiting psychiatric nurse, assuming one is available. Varying levels of care are also implemented in the hospital environment. A patient may have the usual care of routine monitoring or may have more intense one-to-one monitoring if he/she is an imminent risk to him/herself or others.

As a general rule, the more the physical and psychiatric impairments and fewer psychosocial resources, the greater the need to hospitalize for accurate diagnosis and effective treatment. When deciding safe and effective management, the following factors favor hospitalization: poor or unstable physical health, high suicide risk, impaired judgment and reality testing, likelihood of poor compliance, impaired cognitive functioning, lack of social support, and severe anorexia and weight loss.

CO-MORBIDITY OF PHYSICAL ILLNESS: THE INTERFACE OF PRIMARY CARE AND PSYCHIATRY

Accurate diagnosis is a prerequisite for effective treatment. Elderly patients with depression present to their primary care physicians and psychiatrists in a complex manner, and signs and symptoms of physical illness and depression overlap. Even the normal effects of aging may cause diagnostic difficulties and restrict treatment options. Many primary care physicians diagnose and treat late-life depression without referral. However, those patients who fail two or three trials with antidepressants, usually selective serotonin-reuptake inhibitors (SSRIs) or newer agents, are commonly referred to a psychiatrist for further management. These patients represent a treatment challenge and may require complex medication regimens that are more successful with hospitalization. Primary care

physicians also refer for the following reasons: suicidality, co-morbidity with substance abuse, dementia, anxiety disorder, presence of psychosis (delusions, hallucinations), catatonia, bipolar disorder, and inability to tolerate antidepressant treatment^{2,3}. Such patients often need to be managed in the hospital.

Depression is often co-morbid with other physical diseases. Approximately 80% of older adults suffer from at least one chronic health problem⁴. The prevalence of co-morbid depression may be up to 30% in stroke patients, 18% in myocardial infarction patients, 51% in patients with hip fracture, and 50% in patients with chronic pain¹. Existence of an undiagnosed and untreated depression with these illnesses leads to higher disability⁵. The diagnosis of depression with certain illnesses is complex, and the hospital environment provides the necessary monitoring and support staff when complicated medication changes are required.

For example, a patient with cardiovascular disease may present with decreased energy and apathy. Determining whether this is caused by a compromised cardiac status, a medication side effect, or is actually a symptom of depression may be difficult without hospitalization, close monitoring and various medication trials. Formerly, hospitalization was favored for the initiation of tricyclic antidepressant therapy in elderly patients with unstable cardiac disease. First-line treatment with SSRIs is now available and proved safe for use in cardiac disease⁶.

Co-morbid neurological illness is also common in geriatric depression. Patients with depressive symptoms following a cerebrovascular accident also present a diagnostic challenge. There may be communication difficulties or other neurologic abnormalities. Depression may be diagnosed only by the report of the nursing staff and family, who observe apathy, irritability, tearfulness and weight loss⁷. Patients with Parkinson's disease may develop an affective illness or psychosis, which may be secondary to treatment with L-dopa. Hospitalization may be required for medication changes if outpatient support is inadequate.

Severe anorexia, weight loss and refusal to eat are indications for hospitalization for safe and effective treatment^{8,1}. Poor oral intake commonly accompanies severe depression, but it may also result from a variety of medical conditions. For example, individuals with active rheumatoid arthritis may experience insomnia, fatigue and poor appetite equally from their physical illness or an associated depression⁹.

SUICIDE RISK AND THE DECISION TO HOSPITALIZE

Suicidality is the most common reason for psychiatric hospitalization. According to Jacobson²⁹, three goals for inpatient treatment are: (a) the preservation of life and safety; (b) the elimination of

suicidal intent and ideation and treatment of underlying disorders; and (c) the improvement of intrapsychic capabilities, personal factors and psychosocial circumstances to facilitate coping after discharge and decreasing risk of the return of suicidality. However, implementation of these treatment plans is predicted on the initial detection of suicidality.

Careful assessment of suicide risk in depressed older adults is thus vital. The elderly are less likely to have made a prior suicide attempt, but they consistently demonstrate a higher rate of completed suicides¹⁰. The ratio of attempted to completed suicides decreases with age from 200:1 in young adulthood to 4:1 in the elderly¹¹. The higher rate is due primarily to the increased frequency of deaths among older, White males. In 1992, persons 65 and older accounted for 13% of the population but almost 20% of suicides. Even though the frequency of suicide has increased among older persons in the USA, the prevalence is not as high as that of other industrialized societies¹⁰.

Recognition of variables such as gender and race may influence management decisions. Risk factors for suicide in late life include increased age, with the highest prevalence of suicide of persons older than 85¹⁰. Also, being male, White, single, separated or divorced, or widowed are risk factors for suicide. Other risk factors implicated in late-life suicide include: a positive psychiatric history (especially depression and alcohol abuse and dependence); physical illness and functional disability (especially diseases of the central nervous system, malignancies, cardiopulmonary conditions, and urogenital diseases in men); previous suicide attempts; psychological factors (i.e. hopelessness); social factors (stressful life events, e.g. bereavement); and biological susceptibility (dysregulation of the hypothalamic–pituitary–adrenal axis or the serotonin system).

Most older people who commit suicide have seen a primary care provider within 30 days of death¹¹. This observation stresses the need for collaborative efforts with primary care physicians and the need to make careful assessment based on risk factors.

A number of assessment guidelines have been developed to aid in the evaluation of potentially suicidal patients. A four-item screen for identification of suicidal ideation among general medical patients was developed by Cooper-Patrick *et al.*¹².

1. Have you ever felt that life is not worth living?
2. Have you ever thought of hurting or harming yourself?
3. Have you considered specific methods for harming yourself?
4. Have you ever made a suicide attempt?

This four-fold layered approach to assessment is useful in obtaining the necessary data without disrupting the therapeutic relationship. If the answer to the first or second question is negative, then the inquiries can cease and the older person may be considered at low risk for suicide¹⁰. This approach has advantages over other assessment tools, which usually suggest one question to be asked to assess suicidal risk.

DELUSIONS AND LATE-LIFE DEPRESSION

Accurate diagnosis and effective treatment of depressed elderly patients with delusions can be hindered by their impairment of reality testing. Their sometimes well-organized and complex delusions may make them distrust medicine and the physician who prescribes it. This disorder is less frequent in the community and more prevalent in the hospital setting^{13,14}. Accurate diagnosis is necessary, as some studies have suggested that the depression is more severe^{14,15} and it has been associated with suicide¹³. Varying reports also demonstrate decreased cognitive functioning and social functioning among patients with delusional depression¹⁶. These patients are best treated in the hospital. They cannot be relied upon to take accurate doses of medications. Effective

pharmacologic treatment for delusional depression requires combination treatment with high-dose antipsychotic medication and antidepressants¹⁵. ECT has been successful for the treatment of delusional depression^{17–19} and can be performed on an outpatient basis only with adequate social support.

COGNITIVE DYSFUNCTION AND LATE-LIFE DEPRESSION

A full discussion of how depression is distinguished from dementia is given elsewhere in this book. To summarize, diagnosis is difficult because several symptoms of depression and dementia overlap, such as a flattened affect, psychomotor retardation and presence, at times, of delusions²⁰. Delusions are reported to occur in up to 40%¹³ of Alzheimer's disease patients, although they are described as transient and less organized than in delusional depression¹³. Major depression occurs in over 20% of patients with Alzheimer's disease and vascular dementia^{4,21}. This significant co-morbidity may lead to profound disability²².

The diagnosis of depression in dementia usually requires the input of family members or nursing personnel^{23,24}. The patient with cognitive dysfunction and impaired reality testing cannot reliably report symptoms, take medication accurately, or reliably report side effects.

Dementia with depression and behavioral disturbance is frequently too complex to treat on an outpatient basis. These patients may even require involuntary commitment. Aggression may be verbal or physical. Aggression and agitation in dementia may be as high as 50% in the outpatient population^{25,26}. The hospital environment is the only setting with constant monitoring to make an accurate diagnosis, contain behavior and monitor medication.

The reversible cognitive impairment that may accompany depression also increases disability²⁷. With treatment, the cognitive impairment usually improves. However, these patients are at higher risk to develop an irreversible dementia in the future²⁸.

CONCLUSIONS

The patient with late-life depression frequently presents in a complex manner that may require hospitalization. Accurate diagnosis and treatment of these patients is essential to prevent disability, caregiver burden and nursing home placement. Confusion, suicidality and aggression represent psychiatric emergencies in the elderly and may require hospitalization for effective management.

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Electroconvulsive Therapy (ECT)

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Those who decry it in the elderly are sentimental and ill-informed. ECT for suitable patients not only relieves intolerable anguish but saves lives (Brice Pitt¹).

Despite continued opposition to its use, ECT remains a fundamental tool in the armamentarium of the psychiatrist treating the severely mentally ill. There have now been thousands of papers and articles published concerning its use and we have a wealth of accumulated wisdom, and yet the controversy continues².

In geriatric psychiatry there is no controversy: papers continue to confirm that in the elderly ECT is an effective treatment for severe affective disorders³⁻⁶ and is the treatment of choice in severe delusional depression⁷⁻¹⁰. It is safe, despite the likelihood of multiple system disorders and medications¹¹. It is well tolerated, if not always well liked¹², but does sometimes exacerbate confusion. There is no convincing evidence of brain damage or even of lasting memory impairment, particularly if brief pulse right unilateral ECT is used¹³. ECT should be considered in every patient who has either failed to respond to other treatments, or who is suffering intolerable distress, or who may die through inanition or dehydration as a result of his/her depression. It is probably for these reasons that ECT continues to be used six to seven times more frequently in the elderly than in their younger counterparts¹⁴. The fact that in the USA it has also been shown that early use of ECT reduces inpatient costs may have also been an influential factor in its greater use¹⁵.

INDICATIONS

Studies on the efficacy of ECT, which have largely been conducted on younger patients⁷, all emphasize that ECT appears to be more effective than placebo, single-drug therapy and tricyclic/neuroleptic combinations, and that patients with more florid symptoms of recent onset fare best^{7,16-18}. Experience with the elderly would confirm these findings. Indicators of response are perhaps less clear in the elderly, with some authors finding psychomotor disturbance and psychosis a positive predictor of response¹⁰ and others suggesting that patients without these features can also do well¹⁹. That the classic distinction between neurotic and psychotic depression appears less helpful in this group as a predictor of good response is nothing new. Post²⁰ in 1976 stressed the practical irrelevance of any subclassification of elderly depressives, as he found that ECT rendered severe psychotic depressives fit for discharge only slightly more often than neurotic depressives. The presenting picture of hysterical illness, hypochondriasis or other apparent

neurotic illness may well be caused by an underlying functional psychosis, often depressive in the elderly. More recently, in a study of 163 elderly patients given ECT, it was found that 27% had predominantly neurotic depressive features and yet had a good response to treatment³. In fact, in the study by Fraser and Glass²¹, psychic anxiety, along with the more expected features of short duration, severity of illness, guilt and agitation, was one of the symptoms correlated with a favourable response to ECT, whereas the typical endogenous features of late insomnia and diurnal variation in mood were not. Treatment resistance in the elderly will respond to ECT, although not as well as non-resistant patients²². However, they do respond better to ECT than SSRIs¹⁸ and Flint⁵ found that ECT was significantly superior to tricyclic/neuroleptic combinations, even when the former had been augmented with lithium. A pragmatic consensus would suggest that in the elderly a trial of ECT is indicated in any depressed patient who might otherwise be regarded as a treatment non-responder, and if the illness is severe and of short duration there is likely to be a good response, regardless of the presenting symptoms. If the illness has been present for some time, or even years, it still may respond if there is a clear history of a change for the worse in the patient who had previously maintained a stable personality and had coped normally with the vicissitudes of life. Prolonged or abnormal bereavement reactions with marked depressive features not responding to antidepressants or talking therapies may need to be treated with ECT before psychotherapy or counselling can be effective. ECT may need to be given within a few months of the loss if hopelessness and suicidal ideation suggest a risk to life, and should not be withheld due to the feeling that the patient must work through his/her grief naturally, as he/she may never get that chance.

Mania is another indication for a trial of ECT²³, particularly in the elderly, where neuroleptics may fail to control the symptoms and yet produce unsteadiness, postural hypotension or falls, and lithium may not be tolerated due to toxicity problems.

Paraphrenia will often respond to ECT²⁴, although this is more likely if there are obvious depressive symptoms or delusions.

ECT is certainly useful in the depressed patient with Parkinson's disease, as the motor symptoms will improve as well as the depression, and indeed, some authors advocate ECT as the treatment of choice for certain stages of Parkinson's disease, whether or not depression is a major problem^{25,26}. I have given daily ECT with excellent results to a parkinsonian patient who had developed severe paranoid delusions. His refusal to accept his medication rendered him rigid and immobile with pressure sores, he needed intravenous fluids and nasogastric feeding until he had four treatments, whereupon his physical and emotional improvement was dramatic.

Fogel¹⁶ suggests that ECT might be more readily used in the elderly if we were more objective about its virtues as compared with the severe side effects often associated with neuroleptics, which are quite readily used in the agitated elderly patient. Extrapyrarnidal effects were usually the limiting factor but are not so noticeable with atypical neuroleptics. However, he postulates that the demented patient who is very agitated and screaming might suffer less indignity and fewer side effects if treated with ECT, rather than tranquillizers, as the patient may have an underlying affective disturbance manifest only by the agitation and negativism that one often sees in this condition.

CONTRAINDICATIONS

There are no absolute contraindications, only relative risks, relative that is to the morbidity and mortality of untreated depression. The limiting factor is whether the patient is fit for the light anaesthetic that ECT requires. The majority of risk factors are therefore associated with the cardiovascular system. Many people are denied treatment due to irrational caution. For example, pacemakers are not barriers to treatment; the bodily tissues, being highly resistant, prevent the ECT stimulus from reaching the pacemaker in any case. The patient should remain insulated from the ground, however, to prevent the unlikely event of the current leaking to earth and being conducted down the pacemaker wire to the heart. Equally, myocardial infarction is not a contraindication to treatment if the depression is so severe as to threaten life; in less severe cases, an interval, governed by sentiment rather than science, of 4–6 weeks is usually left. The risks are greatest during the first 10 days post-infarct, and probably negligible after 3 months. I have treated a patient with treated hypothyroidism who had two prosthetic heart valves, was therefore on anticoagulants and had a pacemaker, with no special precautions or untoward effects. Patients with osteoporosis or with recent femoral neck fractures can be treated, provided an adequate muscle relaxant is given. Stroke is certainly not a contraindication, and ECT given as soon as 1 month after does not present a major risk to patients. There is now a growing body of literature attesting to the usefulness of ECT in treating post-stroke depression²⁷.

The case of deep venous thrombosis (DVT) is less clear. I have given ECT to a patient who had a DVT in his calf during his depressive illness, once he was adequately anticoagulated. In fact, the risk of pulmonary embolism seems, in my practice, greater in the dehydrated immobile depressive than in those receiving ECT. Arterial hypertension is often regarded as a contraindication, as blood pressure is well known to rise during ECT. This can sometimes be controlled and the pressor response avoided by using sublingual nifedipine or short-acting β -blockers shortly before treatment²⁸. Chronic glaucoma is another condition in which ECT causes fewer problems than tricyclic antidepressants; in fact, intraocular pressures are said to reduce ECT²⁹. Insulin-dependent diabetes is a condition, like Parkinson's disease, which alters during ECT. Insulin requirements may decrease quite substantially during the course of ECT, so more careful monitoring of blood glucose levels is needed. It is also necessary to avoid hyperglycaemia prior to treatment, which may significantly raise the fit threshold, and the timing of ECT administration may need consideration to prevent undue fluctuations in diabetic control. Transient asystole occasionally occurs, for some reason less frequently in the old-old, but it is not of any consequence and need not prevent further treatments^{30,31}.

Epileptic patients on anticonvulsants should not stop their medication during ECT, as that might increase the risk of status epilepticus. However, they may need higher than usual electrical dosages to produce an adequate response.

It is interesting that the seizure during ECT is invested with great powers of harm compared with epileptic seizures *per se*, which can of course occur in patients with any disease or at any time and seldom result in death. It seems understandable, then, that a seizure in the controlled conditions of the ECT room is probably even less likely to result in fatality. There is, of course, a mortality rate associated with ECT but, as noted by Fink³², the treatment rate of 0.002% compares favourably with the rate for anaesthetic induction alone (0.003–0.04%).

ADMINISTRATION

The responses of senior psychiatrists to the process of ECT vary from those who simply prescribe six treatments and leave the administration to the newest recruit, who has often had no training at all, to those surgeons *manqués* who may overstate the risks and precautions in order to increase the perceived risk of their jobs. Clearly, the ideal path lies somewhere between, but nearer the latter than the former! ECT is the only psychiatric treatment in the elderly that involves significant medical intervention with general anaesthesia, and as such, the psychiatrist should have a clear understanding of what he/she is prescribing and regular involvement in its administration. As much attention should be paid to the prescription of ECT as to any other prescription.

A clear decision as to whether bilateral or unilateral electrode placement is wanted should be made; the ECT record sheet should be reviewed to ensure that an adequate convulsive response has occurred without excessive stimulus; treatments should not be in blocks of six, but, provided that the illness is one with a good prognosis, treatment should be continued until the expected degree of improvement is obtained, whether that is after three or 23 treatments.

There is no evidence that the habit of giving one or two extra ECTs after full recovery is effective in preventing relapse³³. The decision to give unilateral or bilateral ECT in the elderly is made easier by the fact that high dose unilateral ECT does seem to produce less confusion, memory loss and headache and appears to be equally effective in many patients^{34,52}. However, there is a great deal of discrepancy in the results of comparative studies, possibly due to differences in diagnosis, age and gender, together with variance in the technique of administration of unilateral ECT. The consensus seems to indicate that, for many patients, both treatments are equally effective; some patients require more right unilateral treatments than bilateral to achieve the same result and some patients who do not respond to right unilateral ECT will respond when switched to bilateral treatment. Male gender and older age are also associated with better response to bilateral treatment.

It is my practice to use bilateral treatment initially in very severe psychotic depressives but right unilateral treatment in most other cases, particularly if there is evidence of prior cognitive impairment, switching to bilateral treatment if there is no response after six to eight right unilateral treatments. Brief pulse ECT at a moderately supra-threshold stimulus (which is often only around 275–350 millicoulombs) appears to offer efficacy, with the advantage of much less memory loss and confusion than the modified sine wave stimulus, and should be used in all cases, with a record of dosage received by the patient to ensure adequate technique⁵³.

There is considerable debate about the necessity to use a dose-titration technique to establish seizure threshold prior to treatment, with some viewing this as unnecessary and even detrimental in those patients requiring several non-convulsive stimuli. Adequate seizure response can be measured using inter-ictal EEG monitoring. Some clinicians seem to have developed an over-weaning

interest in stimulus intensity, seizure threshold and seizure duration. It is clear that seizure threshold will increase by about 40% during the course of ECT and the seizure duration will tend to decrease by about one-third. However, while it seems that outcome is not correlated with either seizure duration or threshold for bilateral ECT³⁵, it may be more important to keep the stimulus intensity above seizure threshold in unilateral treatment³⁶. Seizure duration is more difficult to evaluate, as some patients regularly have brisk and brief responses with good results. The use of propofol to induce anaesthesia consistently reduces seizure duration, although apparently without affecting efficacy^{37,38}. The use of caffeine prior to treatment to prolong seizure activity has been associated with improved efficacy in some patients³⁹. It would seem that, as a rule of thumb, we should aim for a seizure length of around 25 s and any seizure less than 15 s or more than 120 s is likely to adversely affect response⁴⁰. Cumulative seizure duration again seems less interesting now than it once was as a measure of the length of a course of treatment, and clinical response still seems the best measure.

There is no evidence that routine atropine premedication improves cardiac stability or lessens secretions. Theoretically it could cause confusion, but there is no convincing evidence of this either. Glycopyrrolate, which does not cross the blood-brain barrier, may be a better drug to use as a drying agent. Methohexitone for the induction of anaesthesia at a dosage of 30–50 mg is adequate to ensure sleep without hangover, and muscle relaxation with a suxamethonium dosage of 20–40 mg is enough to modify the convulsion without abolishing all evidence of motor activity. If the minimum amount of anaesthetic is combined with treatment early in the morning, the patient is not required to starve any longer than usual, he/she is less likely to be dehydrated, is less likely to break his/her fast, has less time to become anxious and agitated and will recovery quickly enough to enjoy a breakfast with the other patients on the ward. If this routine is combined with regular supervision of treatment by the prescribing psychiatrist, the patient will derive the maximum benefit from each treatment and the course will not be unnecessarily prolonged or ineffective. Outpatient ECT does not appear to be as effective in the elderly, except occasionally as maintenance, and consequently most patients will require admission to a specialist unit, where the effects of ECT combined with the therapeutic milieu will hasten improvement. Familiar staff administering the treatment and a well-designed ECT suite will help reduce anxiety.

MAINTENANCE AND CONTINUATION ECT

In 1990 the American Psychiatric Association task force on ECT defined continued administration of ECT over a 6 month period to prevent relapse after induction of remission as continuation ECT (C-ECT); treatment beyond 6 months was termed maintenance ECT (M-ECT). This was felt to be a viable form of management for selected patients.

Maintenance ECT has been used for many years: a survey of British psychogeriatricians in 1991⁴⁷ found that 20% were using it but there is little more than anecdote to support its use in the literature. Such studies as there are consist mainly of case-studies and small series of hospitalized patients, all of a “naturalistic” nature.

In a 1 year follow-up of nine elderly patients, continuation treatment, even if discontinued fairly quickly seemed to confer some lasting advantage in prevention of relapse⁴⁸, as did Petrides *et al.*'s study, looking at 33 courses of C-ECT⁴⁹. The conclusion seemed to be that where patients have responded to acute ECT but previously failed on continuation pharmacotherapy there was compelling evidence for C-ECT and little therapeutic alternative.

The four patients in this study⁴⁹ who continued with M-ECT remained well and the five who had previously stopped did not. Naturally this result is open to other interpretations, but it does suggest that C-ECT should be considered for those with recurrent depression who respond well to ECT acutely but receive no prophylaxis from pharmacotherapy. The practicalities of using outpatient M-ECT have prevented my using it more. Bringing elderly patients to hospital for outpatient ECT early in the day, from a rural catchment area some distance from the hospital, can be problematic, they soon lose enthusiasm for the treatment and consequently often withdraw consent. This is an issue recently addressed by Kim⁵⁰. However, Schwarz's findings, that rehospitalization rates were reduced by 67% after instituting M-ECT, suggest that we should try and overcome the practical difficulties⁵¹.

CONSENT

Popular myths about ECT are always more readily believed than the reality and can be part of what the patient believes they are consenting to. Occasionally patients consent as part of their death wish. I use a video of myself administering ECT to a patient seen before and after treatment, to show anxious or interested relatives and patients; no-one having seen it has then declined the treatment. There is one study suggesting that understanding is not enhanced by this method. The issue of informed consent in depressed patients is complex. As I have suggested, many care little and are prepared to do anything their doctor suggests, and patients' recollection of what was explained to them, after the ECT and when the depression has lifted, is often vague. A careful explanation should be made and recorded, and if there are doubts on either side a chance to preview the ECT room or an explanatory video may be helpful. However, it is doubtful whether the explanation of ECT is any less detailed than that of most surgical procedures and most people are willing to consent without seeing a video of the operation in question. Passive acceptance of ECT is often the case in the severely depressed but this should not prevent a full explanation, including consulting relatives if appropriate.

Involuntary ECT should never be given except within the guidelines of the relevant Mental Health legislation if we are to ensure the availability of ECT as a treatment option in the future. Nevertheless, depression is such a serious and debilitating illness that the chance of a cure through use of ECT should never be denied to a patient whose prognosis is favourable, simply through difficulty in obtaining actual written consent.

SIDE EFFECTS

As already mentioned, confusion and memory loss are often regarded as an inevitable corollary of ECT in the elderly, but this is clearly not the case and there are well conducted studies showing no objective permanent effects on memory, and in fact this often improves as a result of improvement in the depression^{21,41,42}. Nevertheless, there is no doubt that some patients who were given bilateral sine wave ECT experienced long-term, even permanent, memory loss, and bland reassurances that this or even brief pulse bilateral ECT will not cause any memory loss is foolish and counterproductive. Some patients given bilateral brief pulse ECT may have amnesic gaps, but can be assured that no lasting effect on memory function, i.e. new learning or intelligence, will occur. The situation with right unilateral brief pulse ECT is different, with any subjective memory impairment being transient and undetectable 6 months later^{43,44}. Patients with existing dementia may well show signs of memory impairment, even

with unilateral ECT. This may be acceptable in view of the relief from distress and agitation and improvement in behaviour and performance.

The cognitive side effects can be minimized by reducing concomitant medications, particularly benzodiazepines, anticholinergic antidepressants and lithium⁴⁵, although a recent study found no problems with the administration of ECT and lithium⁴⁶. Benzodiazepines, given intravenously as the seizure ends, can be of use in controlling emergence delirium, which can last for 15–30 min after treatment and be very difficult to control otherwise¹³. Dementia *per se* does not preclude the use of ECT, provided that the coexisting depression is circumscribed. A history of depression before the dementia adds weight to the decision, particularly if there was a good response to ECT previously. Other side effects of treatment, such as headache and dizziness or muscle pain, usually only after the first anaesthetic, are minimal and soon forgotten as the depression lifts.

ECT is a valuable and as yet essential tool in the treatment of depression in old age, a disease which untreated carries a significant mortality. It is interesting that in my practice elderly patients who have attempted suicide are nearly all offered ECT. This is because those, albeit only very few in number, who have subsequently killed themselves during a depressive illness have all been patients who have either refused ECT or not been given it at the time of their index suicide attempt. However, whilst there are many compelling arguments for the use of ECT, it is not a universal panacea. ECT, like any potent treatment, should be prescribed with accuracy and its use monitored carefully by those prescribing it.

Depression in the elderly presents with protean manifestations. ECT should be part of an eclectic approach to treatment and as such will continue to relieve distress and save lives.

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Pharmacological Treatment of Depression

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A recent review¹ identified mood disorders as a major public health problem, with poor recognition, diagnosis and treatment despite the availability of reasonably safe, effective, economical treatments and the established effectiveness of continuing educational programmes for care providers. The magnitude of the following problems is greater in the elderly with mood disorders: poor recognition for most somatic and cognitive symptoms; increased physical morbidity and disability; and high mortality from suicide and other causes. This situation, however, is balanced by the availability of effective pharmacological treatments, which are successful in two-thirds of patients within few weeks of treatment, who remain well 1 year later. However, the temporal profiles of the course of late-life depression during treatment shows marked variation in rate, stability and direction of recovery with reliable pretreatment predictors of outcome². The usefulness of pharmacological treatment for patients with subsyndromal depression has not, however, been investigated³.

PRETREATMENT CONSIDERATIONS

There are a number of pretreatment considerations before starting specific pharmacological treatment of mood disorders in the elderly^{4,5}. With advancing age, there are important and clinically significant changes in distribution, metabolism and elimination of these drugs⁶. The age-related increase in volume of distribution results in a longer half-life for all psychotropic drugs. Hepatic drug metabolism decreases with age, also resulting in a prolonged half-life, which may be two to three times longer than in younger patients, and the decrease in renal function with advancing age is particularly relevant with regard to lithium, which also results in two to three times higher plasma levels than those in younger patients on the same daily dose. Of particular importance, however, has been the study of the inhibitory effects of the selective serotonin reuptake inhibitors (SSRIs) on the cytochrome p450 enzymes; these pharmacodynamic actions have pharmacokinetic consequences for co-administered drugs, such as tricyclic antidepressants (TCAs), which are dependent on these enzymes for biotransformation.

To determine the best pharmacological treatment options for individual patients requires careful consideration of a number of clinical factors, which include the following: type of mood disorder; degree of urgency for treatment; previous response to treatment; concurrent medical problems; concurrent drug therapy; risk of overdose; reasonable half-life; dosing flexibility, and affordability⁷.

TREATMENT OF DEPRESSION

Elderly patients with depression could be successfully treated with conventional TCAs, monoamine oxidase inhibitors (MAOIs), SSRIs and atypical antidepressants. A recent update of the evidence reported the results of 26 RCTs of pharmacological treatments and 21 of these were placebo-controlled. People were recruited mainly from outpatient clinics. Significant benefits were found for fluoxetine, trazodone and phenelzine. Of the 17 drug vs. drug comparisons (mainly involving heterocyclic drugs), none showed significant benefit above the others. In people with depression plus a physical illness vs. placebo, one systematic review of 18 RCTs found that antidepressants were more effective than placebo in people with depression and a physical illness. TCAs have the major limitations of anticholinergic effects, postural hypotension, excessive daytime sedation and cardiotoxicity in overdose. Their advantages, however, are their established efficacy and low cost. A number of recent reviews^{4,6,7} identified the secondary amines desipramine and nortriptyline among the TCAs having more favourable side-effect profiles, with desipramine having the fewest anticholinergic effects and nortriptyline causing the least postural hypotension. The secondary amines also have the advantage of therapeutic drug monitoring, with an established therapeutic window for nortriptyline and a therapeutic plasma level of desipramine. Therapeutic drug monitoring enables the clinician to determine the minimal therapeutic dose and to monitor compliance. Nortriptyline has been well investigated for use in elderly patients, including patients older than 80 years⁸ and patients who have had strokes⁹.

Among the MAOIs, phenelzine has been shown to be effective in treating elderly patients with depression^{10,11}, particularly for patients who could not tolerate TCAs and those with resistant depression. Their main limitation is their interaction with tyramine-rich foods, causing hypertensive crisis. For this, they are superseded by moclobemide, a reversible and selective inhibitor of monoamine oxidase type A. Comparative trials in the elderly have established its efficacy compared with imipramine¹², nortriptyline¹³ and placebo¹⁴. Moclobemide also showed enhancing effects on cognition in patients who had dementia and depressive symptoms¹⁴.

The SSRIs, however, have provided a major advance in the successful and safe management of depression in late life. These include fluoxetine, fluvoxamine, sertraline, paroxetine and citalopram. The advantages of SSRIs over TCAs are the absence of anticholinergic effects, orthostatic hypotension and arrhythmia in the side-effect profiles, and their safety in overdose⁷. The consensus statement update from the US National Institutes of Health³ on the diagnosis and treatment of depression in late life concluded that the SSRIs are roughly equivalent to TCAs in

efficacy in the elderly, with 60–80% of patients responding to treatment. The authors concluded that the efficacy data are not robust when an SSRI is compared with placebo; the only available report from a randomized placebo-controlled trial of fluoxetine in outpatients¹⁵ found a lower drug–placebo difference than that found for many studies with TCAs. Moreover, comparison of fluoxetine with nortriptyline in inpatients with severe depression and heart disease showed that nortriptyline may be more effective in this population¹⁶. A meta-analysis of the comparative efficacy and safety of SSRIs and TCAs in elderly patients¹⁷ showed no differences in safety and dropout rates.

The side-effects profile of SSRIs includes nausea, diarrhoea, insomnia, headaches, agitation, anxiety and sexual dysfunction. The SSRIs also seem to worsen parkinsonism. As with other antidepressants¹⁸ there have been case reports of hyponatraemia, hypomania and seizures⁷. An important aspect of their use is their drug–drug interactions, and their inhibitory effects on hepatic cytochrome p450 isoenzymes, the route through which many drugs commonly prescribed for elderly people are metabolized¹⁹. Paroxetine, norfluoxetine and sertraline have clinically important inhibitory effects (*in vivo*) on cytochrome P2D6, resulting in increased plasma concentrations of co-administered TCAs, such as desipramine, and antipsychotics, such as haloperidol²⁰. Sertraline, however, had a modest effect on plasma nortriptyline levels in depressed elderly patients²¹. Fluoxetine increases plasma levels of co-administered carbamazepine; alprazolam and fluvoxamine increase plasma concentration of co-administered TCAs and antipsychotics by inhibiting cytochrome P1A2⁹. An extensive list of drugs metabolized by various p450 isoenzyme types is provided in the reviews by Catterson *et al.*⁵ and Rivard⁷.

CONTINUATION AND PROPHYLACTIC TREATMENT

Early studies indicated a poor outcome of late-life depression²⁵, with high relapse, recurrence and chronicity. This view was based on naturalistic observation without monitoring of compliance, adequate dosage and duration of treatment, including prophylactic treatment, and was therefore challenged²⁶. Controlled studies of maintenance antidepressant medication, however, showed better outcome for late-life depression²⁷. The Pittsburgh group reported a 3 year follow-up study of maintenance treatment with nortriptyline or placebo with or without interpersonal psychotherapy (IPT)²⁸ and showed that 80% of patients assigned to nortriptyline, with or without IPT, remained in remission. The Pittsburgh group also identified the elderly patients who remained well after placebo-controlled discontinuation of antidepressant medication for a period of 1 year. Recovery of good subjective sleep quality by early continuation treatment was useful in identifying which remitted elderly depressed patients remained well with monthly IPT after discontinuation of antidepressant medication²⁹. Moreover, effective maintenance treatment with nortriptyline was associated with enhancement in the rate of delta-wave production in the first non-rapid eye movement sleep and of rapid movement activity throughout the night³⁰.

A recent study of the effect of treatment on the 2 year course of late-life depression³¹ showed a 74% survival rate without relapse. This good outcome was obtained by the use of full-dose antidepressant medication, frequent follow-up and rigorous treatment of relapse.

The US National Institute of Mental Health consensus statement update³ concluded that recent evidence supports the recommendation for at least 6 months of treatment beyond recovery for those with first onset in late life, and for at least 12 months for those with a recurrent illness³². Moreover, prophylactic treatment should be of the same type and of same dosage as

that which was successful in the initial acute phase. The consensus statement also concluded that treatment response and long-term outcome for all the patients is generally similar to that observed in younger adults, but the temporal course may be somewhat slower in the elderly and risk of relapse somewhat greater³. The use of lithium in continuation and prophylactic treatment of depression in the elderly has also been recommended^{27,33,34}, with the use of lower doses/plasma levels³⁴. Lithium may be a more favourable prophylactic treatment than antidepressants in recurrent depression with melancholia and in depression with psychotic features (delusional depression), which are particularly common among the elderly, with a tendency to respond less well to antidepressants^{26,34}. The other advantage of lithium therapy is the evident decreased mortality, whether from suicide or other causes³⁵. A recent 1 year prospective, placebo-controlled study of maintenance lithium in conjunction with cognitive–behavioural psychotherapy in elderly depressed patients³⁶ showed that, although cognitive–behavioural psychotherapy reduced depression severity during follow-up, lithium therapy was no better than placebo. This appears to be related to poor compliance, a finding that highlights the serious difficulties in undertaking prophylactic studies in elderly depressed patients.

TREATMENT OF BIPOLAR DISORDER

Elderly patients with late presentation or late-onset mania respond well to standard antimanic treatment with neuroleptics, lithium and anticonvulsants³⁷. Neuroleptic treatment is best avoided in the elderly because of its known extrapyramidal side effects, except for floridly psychotic, agitated and behaviourally disturbed patients who need rapid control of symptoms. Lithium remains the treatment of choice, followed by valproic acid³⁷. The evidence for the efficacy of lithium in late-life mania is based on retrospective and uncontrolled studies; there have been no controlled studies, and there have been no controlled studies of the efficacy of anticonvulsants in late-life mania. It has been suggested that valproate is a safer alternative treatment to lithium than carbamazepine, whether used as single or adjunct treatment, in elderly manic patients³⁷. There are no guidelines regarding the optimal plasma concentration of valproate in relation to efficacy³⁷.

A recent evidence-based review of the treatment of mania, mixed state and rapid cycling in younger populations³⁸ concluded that lithium and divalproex sodium are effective in mania, whereas divalproex sodium and carbamazepine are more effective in mixed states. Divalproex sodium is the drug of choice for rapid cycling disorder. With bipolar depression, lithium is recommended as a first-line treatment and the addition of a second mood stabilizer or the TCA would be an appropriate next step³⁹.

The guidelines for the continuation and prophylactic treatment of bipolar illness in late life are similar to those advocated for younger patients, except for the notion of high recurrence rates necessitating prophylactic treatment, even after a first-onset manic episode. Lithium remains the medication of choice for prophylaxis³⁴. An open naturalistic study of lower doses/plasma-lithium levels of lithium³⁴ showed efficacy in the elderly comparable to that in younger patients at plasma lithium levels as low as 0.4 mmol/l, with fewer side effects and renal and thyroid adverse effects.

TREATMENT-RESISTANT DEPRESSION

Treatment-resistant depression occurs in one-third of elderly depressed patients⁴⁰ and can only be ascertained after adequate recognition, compliance with treatment and effective treatment⁴¹.

It has also been related to cognitive impairment, physical and psychiatric co-morbidity, late onset and presence of melancholic and psychotic features⁴¹. Moreover, elderly patients with anxious depression are less responsive to nortriptyline than are those without significant anxiety symptoms⁴².

Although TCAs¹⁶ and MAOIs⁴³ have shown efficacy, the SSRIs are specifically advocated⁴¹. Fluvoxamine has shown efficacy (70% good response) in desipramine non-responders^{44,45}, and patients who were intolerant of fluoxetine completed a trial of sertraline with a response rate of 76%⁴⁶. Efficacy has also been shown for trazodone⁴⁷, bupropion⁴⁸ and venlafaxine⁴⁹. Combination and augmentation strategies have been advocated⁴¹. Lithium augmentation in TCA non-responders is effective in 20–65% of cases^{50–52}. It is, however, conducive to cognitive and neurological side effects in 50% of patients^{50,51,53,54}. Lithium has been successfully added to SSRIs, notwithstanding the risk of neurotoxicity with an SSRI–lithium combination⁴¹. Advocated augmentation/combination strategies includes TCA/triiodo-thyronine; SSRI/TCA; SSRI/anticonvulsants; and SSRI/oestrogen⁴¹, and elderly patients requiring adjunctive medication to achieve remission may need continuation of adjunctive medication to remain well and to avoid early relapse⁵⁵.

For refractory bipolar disorders, a recent review⁵⁶ concluded that the safest combination of mood stabilizers is valproate plus lithium. This was also shown in a series of elderly patients with lithium-resistant rapid cycling mania⁵⁷.

CONCLUSION

Although there have been impressive advances in the pharmacological treatment of mood disorders in general, there has been a relative paucity of controlled studies in the elderly, particularly in maintenance and prophylaxis. Generalization from the results of studies of younger patients may be inappropriate in view of the significant changes associated with normal ageing and concomitant medical illness, which affect the pharmacokinetics and pharmacodynamics of psychotropic drugs.

Nevertheless, there has been a change of culture. The nihilism that had prevailed in the treatment of mood disorders in late life has been replaced by cautious optimism with regard to the results of controlled trials in naturalistic settings, as well as studies in high-risk groups, including patients with multiple medical conditions and subsyndromal states.

A large majority of elderly patients with depression could be treated successfully with antidepressants, particularly the SSRIs, because of their favourable side-effect profiles and their low toxicity in overdose. The SSRIs, however, challenge the clinician with their clinically significant drug–drug interactions. Patients who improve should receive continuation of prophylactic treatment with the same dose. For mania, lithium remains the optimal treatment, with anticonvulsants, particularly divalproex, providing a second-line treatment. The efficacy and safety of atypical neuroleptics remain to be evaluated in both acute and long-term management of bipolar illness. There is also hope for those with resistant-mood disorders with the design of augmentation/combination strategies, which require further evaluation.

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Treatment-resistant Depression

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Treatment resistance is a common clinical problem, reported in up to one-third of older depressed patients¹. Many patients labelled as “treatment-resistant” in fact have not had an adequate course of treatment². Therefore, the first step in achieving remission of depressive symptoms is to ensure that the patient has been given, and has complied with, an optimum dose of antidepressant for a sufficient length of time (at least 6 weeks). In treatment-resistant depression, it is also important to investigate the patient for unidentified physical conditions (such as hypothyroidism, vitamin B₁₂ or folate deficiency, or hypercalcemia) that could be contributing to poor antidepressant response².

In patients who have failed to respond to an adequate trial of antidepressant medication, the following options can be considered: (a) augment the antidepressant with another drug that is not primarily an antidepressant, such as lithium, triiodothyronine, methylphenidate, pindolol, buspirone or valproate; (b) add a second antidepressant to the first (combination therapy), e.g. add

a tricyclic antidepressant (TCA) or bupropion to a selective serotonin-reuptake inhibitor (SSRI); (c) switch to a different antidepressant medication; or (d) switch to electroconvulsive therapy (ECT). The advantage of augmentation or combination therapy is that they do not require discontinuation of the original antidepressant and, therefore, patients who have partially responded to treatment are not put at risk of returning to their baseline severity of depression. Also, response may at times be faster with augmentation/combination than with a new trial of antidepressant medication. The disadvantage of these strategies, especially in older people, is that the combination of medications increases the risk of side effects and drug–drug interactions. Also, there have been no placebo-controlled trials of augmentation or combination therapy in elderly depressed patients and so their efficacy has not yet been established in this population^{2,3}.

There are virtually no research data on switching from one antidepressant medication to another in refractory geriatric

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In patients who have failed to respond to an adequate trial of antidepressant medication, the following options can be considered: (a) augment the antidepressant with another drug that is not primarily an antidepressant, such as lithium, triiodothyronine, methylphenidate, pindolol, buspirone or valproate; (b) add a second antidepressant to the first (combination therapy), e.g. add

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There are virtually no research data on switching from one antidepressant medication to another in refractory geriatric

depression. Flint and Rifat⁴ found that seven of 15 elderly patients (47%) who had failed to respond to 6 weeks of nortriptyline followed by 2 weeks of lithium augmentation subsequently responded to a 6 week trial of phenelzine. This rate of response is comparable to that reported among younger patients refractory to TCAs⁵. Data obtained from mixed-aged patients with resistant depression suggest that switching to an antidepressant within the same class is less effective than switching to one from another class⁵. If a patient has failed to respond to an SSRI or nefazodone, then switching to an antidepressant with dual neurotransmitter action (e.g. venlafaxine, mirtazapine or nortriptyline) is a reasonable approach.

Although substitution of treatment usually involves switching from one antidepressant medication to another, switching from an antidepressant to ECT should also be considered. The efficacy of ECT in patients who have failed to respond to adequate antidepressant pharmacotherapy is lower than in patients without established medication resistance⁶. Nevertheless, no antidepressant, alone or in combination, has been shown to be more effective

than ECT in treatment-resistant depression and ECT remains a valuable option in this situation.

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Psychotherapy of Depression and Dysthymia

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BACKGROUND

Depression in the elderly is a treatable disorder. Although antidepressant medication has traditionally been considered a first-line intervention, psychotherapy has been shown to be *at least* equally effective¹⁻³. The NIH Consensus Development Conference on Diagnosis and Treatment of Depression in Late Life concluded that psychotherapy was an important treatment option for elderly depression, although not sufficient by itself⁴⁻⁶. This conclusion has been challenged in the literature, with reviews of psychotherapy research finding a powerful effect of psychotherapy alone on depression as compared to wait-list controls^{1,3,7}. Additional research on the combination of medication and psychotherapy for elderly depression has shown promising results for both acute treatment and relapse prevention⁸. Less is known about the treatment of dysthymic disorder, although its similarities to major depressive disorder suggest that aggressive treatment is the most powerful treatment option.

This chapter reviews the theoretical elements and empirical evidence supporting the use of psychotherapy to treat elderly depression. First, we examine special issues associated with treating elderly depression using psychotherapy. Next, we review the theoretical foundations and evidence for the use of psychotherapy with older adults, including cognitive-behavioral, psychodynamic, interpersonal and group interventions. Finally, we examine recent theoretical developments and research that have addressed problems associated with relapse prevention, treatment-resistant depression, and co-morbid personality disorders.

SPECIAL ISSUES IN PSYCHOTHERAPY WITH OLDER PATIENTS

Psychotherapy is a particularly useful option for depressed patients who cannot or will not tolerate medication, or who are dealing with stressful conditions, interpersonal difficulties, limited levels of social support and/or recurrent episodes of depression^{6,8,9}. Although it has been established that psychotherapy and medication are effective treatments for major depression, there is a sizeable literature highlighting the underdiagnosis and undertreatment of late-life depression¹⁰⁻¹⁶. Consequently, millions of older persons deserving of mental health care go untreated¹⁷. Friedhoff¹⁴ indicated that only 10% of older adults in need of psychiatric services actually receive professional care and there

has been minimal utilization of mental health services in this age group^{16,18,19}.

Under Diagnosis

Older adults are more likely to see general practitioners than mental health professionals, essentially leaving the responsibility for diagnosis of late-life depression in primary care. Although physicians are more likely than mental health professionals to see depressed older adults, this does not mean that elderly patients will purposefully present with depressive symptoms. Studies have shown that up to 75% of older adults who complete suicide had seen their physician in the previous month, emphasizing the important role general practitioners play in diagnosing clinical depression²⁰. In a non-depressed community sample of 462 older adults, 40% indicated they would keep depressive symptoms to themselves and not report them to anyone²¹. Of those who would seek help, over half (52%) indicated they would talk to their regular doctor. Allen *et al.*²² surveyed attitudes toward treatments for depression in older and younger inpatients. The older group, particularly those identified as depressed, was less likely to approach anyone for help. In addition, the older adults had more negative attitudes about approaching their physician than the younger group.

Hesitation to seek help for mental health problems should not be considered unfounded. Link and colleagues²³ have identified a social stigma associated with undergoing treatment for psychiatric dysfunction. However, treatment outcome research suggests that gains in well-being likely outweigh any stigma associated with the disorder. Hence, coming to an understanding of the origins of these attitudes may assist in developing educational interventions addressing the needs of this age group.

This descriptive research highlights several issues of concern for the diagnosis of depression. First, general practitioners typically bear the responsibility for the diagnosis of depression in their older patients, despite the possibility that patients may not feel comfortable about discussing depressive symptoms with others. Making this more difficult have been reports that older patients may not recognize that they have symptoms, or may pass off symptoms as signs of other physical conditions¹⁰. Second, not all older patients regularly see general practitioners. For these older adults, informal resources (family, friends, religious sources) are left with the responsibility of identifying changes in moods and the need for professional intervention. Third, psychiatrists and

psychologists have greater expertise for accurate diagnosis, yet typically only encounter a potential patient after referral from a general practitioner. In addition, mental health professionals specializing in work with geriatric populations are still in the minority and most training programs for both psychology and medicine do not emphasize geriatric mental health issues. This only exacerbates the factors that make recognition less likely to occur^{24,25}. In order for improvements in detection to occur, all three areas discussed above will need to be addressed. Physicians will need to enhance their ability to detect depressive symptoms and make quick treatment decisions during increasingly short office visits. Relatives and other informal sources of referral will require greater education regarding depressive symptoms and encouragement to report those symptoms. Finally, a greater number of mental health professionals will need to obtain specific training in geriatric issues and increase their visibility among potential referral sources.

Under Treatment

Ideally, if the dilemma of underdiagnosis of late-life depression could be resolved, under treatment would not be a problem. However, this does not appear to be the case. Research indicates that even cases that are diagnosed are not necessarily treated. Koenig *et al.*¹³ reported that 34% of hospitalized medically ill elderly patients with a diagnosis of depression in their charts were not treated with antidepressant medication during their inpatient stay or during a 45 week follow-up period, neither were these patients referred to psychotherapy. Only 12.4% of the sample had documented psychotherapeutic intervention in their medical charts at discharge¹³. Thus, not only does recognition appear to be a difficulty, but under-treatment as well. The question is why?

Many practitioners assume that older adults have negative attitudes toward psychotherapy as a treatment for depression. However, research on attitudes toward treatment in elderly samples is not conclusive, with considerable descriptive research suggesting that older adults may prefer counseling over medication treatment for depression. Sixty-eight percent of the previously mentioned non-depressed community-dwelling older adult sample agreed with a statement that professional counseling or therapy helps most depressed people feel better. Interestingly, 56% of the same sample reported that they believed antidepressant medications to be addictive, and only 4% disagreed²¹. Older adults have also been shown to report a greater number of positive attitudes toward mental health professionals, and to be less concerned with stigma attached to seeking treatment for depression relative to younger adults²⁶. On the other hand, Allen *et al.*²² found attitudinal barriers to treatment in a survey of both younger adult and older adult inpatient samples. However, in the older sample, the pattern of preference for counseling and psychotherapy over medication persisted, with 95% of the older group agreeing that "people with depression should be offered counseling", and 46% agreeing that "people with depression should be treated with antidepressant tablets"²². Hence, it appears that interventions addressing these issues must educate practitioners to evaluate their assumptions about elderly preferences for treatment, as well as educate older and younger adults about available treatment options.

PSYCHODYNAMIC THERAPY

This type of treatment is based on psychoanalytic theory that views current interpersonal and emotional experience as influenced by early childhood experience. This experience results

in the development of a complex inner world shaped by both unconscious and conscious mental processes. There have been a variety of theoretical formulations developed over the years that have utilized a psychodynamic formulation to treat depression, including classic psychoanalytic theory²⁷, ego-psychology²⁸, self-psychology²⁹ and object-relations theory³⁰. Revised conceptualizations have emphasized understanding how relationships are internalized and transformed into a sense of self²⁹⁻³². Because early interactions with caregivers are so tied up with emotional gratification and deprivation, the interaction with mother is viewed as a template for all subsequent relationships. Psychopathology is theorized as related to arrests in the development of the self and depression is viewed as a symptom state resulting from unresolved intrapsychic conflict, which may be activated by life events such as loss.

There have been several indications in the geriatric depression literature that short-term psychodynamic therapy, particularly as conducted by Thompson, Gallagher-Thompson and colleagues^{33,34}, is an effective means to treat depression in older samples. In studies with random assignment to wait-list control, short-term psychodynamic therapy or cognitive-behavioral therapy, there were no significant differences between the types of psychotherapy at the end of treatment or at 12 and 24 month follow-ups. Additional research on depressed caregivers demonstrated an interaction between mode of therapy and length of caregiving, such that those who had been providing care for less than 44 months appeared to improve more from dynamic therapy, whereas longer-term caregivers seemed to benefit most from CBT³⁵. The authors suggested that the long-term caregivers needed the skills learned in CBT in order to care for family members with more pronounced deficits and requiring more complicated care. These interesting results call for additional controlled trials comparing different treatment modalities, continued component analysis research, and continued research that examines which type of treatment works best with which type of patient.

LIFE REVIEW AND REMINISCENCE THERAPY

Life review and reminiscence therapies are psychoanalytically orientated approaches to psychotherapy for depression. Life review therapy includes a review of life experiences in order to revisit and resolve old conflicts and reintegrate life experiences^{36,37}. Conflicts emerge in interviews about past life experiences, in both acknowledgment and omission. Reminiscence therapy differs from life review in that the focus is on enhancing self-esteem and social intimacy by recounting past experiences, rather than directly resolving past conflicts^{2,38}. Lewis and Butler³⁷ suggest several techniques to encourage elderly clients to participate in the life review process. Written and taped autobiographies, pilgrimages to the place of childhood, reunions, scrapbooks and photographs are tools that can stimulate past memories. A crucial component of any life review therapy is careful, attentive listening on the part of the therapist^{36,38}.

Results on empirical research of life review and reminiscence therapies are promising, but inconsistent. Teri and McCurry² reviewed 12 empirical studies using reminiscence techniques. Results were mixed, likely due to the range of patient populations (institutionalized, homebound, community-dwelling), treatment duration (1-10 sessions), type of therapy (group, individual) and diagnosis (major depression, no diagnosed psychopathology). Also, the distinction between life review and reminiscence therapy has not been clearly operationally defined, resulting in an ambiguous designation of the differences between the two therapies.

Research does suggest that the amount of structure in reminiscence therapy may influence outcome³⁹. Fry³⁹ compared individual structured reminiscence therapy, non-structured reminiscence and non-reminiscence visits in a sample of moderately depressed elderly. Subjects in the non-reminiscence control group had significantly higher post-treatment depression scores than those in either reminiscence group; the subjects in the structured reminiscence groups had lower depression scores than those in the unstructured reminiscence groups. Reminiscence group therapy has also been tested in nursing home residents with dementia and has been found to reduce self-reported symptoms of depression to a greater degree than subjects in supportive therapy or control groups⁴⁰. However, the therapy did not improve cognitive or behavioral outcomes, and the reduction in depression was short-lived⁴⁰. Rattenbury and Stones⁴¹ found that both reminiscence and current event discussion groups showed positive changes on measures of psychological well-being when compared to non-treatment controls. However, because subjects were not selected for high depression scores, this was not an empirical test of treatment for *clinical depression*. Yet the results do suggest that there is something beneficial about client-focused interaction, a fundamental component of any form of psychotherapy.

INTERPERSONAL PSYCHOTHERAPY

Interpersonal psychotherapy (IPT) is a manualized, time-limited outpatient treatment for depression, focusing on current interpersonal issues in four problem areas: interpersonal disputes, role transitions, interpersonal deficits and abnormal grief⁴². The therapist and client collaborate to identify which problem area to focus on in treatment; commonly, more than one problem area is chosen. Klerman *et al.*⁴² pointed out that, regardless of the origin of depression (genetic, biochemical, developmental vulnerability, personality), the condition is expressed within an interpersonal context. The initial goal of therapy is to reduce symptoms of depression, but the overarching goal is to improve the patient's social functioning and interpersonal relationships⁴³. With its emphasis on addressing interpersonally relevant problems, IPT appears particularly well suited to the life changes that many older people experience. Techniques utilized in treatment include: role playing, communication analysis, clarification of the patient's wants and needs, and links between affect and environmental events⁴⁴. Frank and colleagues⁴⁵ have developed separate treatment manuals for IPT in late life (IPT-LL) and interpersonal maintenance therapy for older patients (IPT-LLM) (cited in ref 44). These manuals include adaptations specific for use in elderly patients, including, but not limited to, flexibility in length of sessions, long-standing role disputes, and the need to help the patient with practical problems.

Controlled trials in adult depressed populations have demonstrated the efficacy of IPT for the treatment of acute depression (reviewed in refs 43, 44). IPT has also been found as effective in the acute treatment of major depressive disorder in elderly patients as nortriptyline⁴⁶. Of additional importance were findings that elderly patients in IPT treatment were less likely to drop out of treatment than those taking nortriptyline, because of the medication's side effects.

Research from the Reynolds group at the University of Pittsburgh has shown IPT in combination with nortriptyline to be an effective treatment for elderly depression in geriatric samples^{47,48}. In an attempt to understand more regarding the treatment of elderly patients with recurrent depression, Reynolds *et al.*⁴⁷ selected patients only if they reported at least one prior episode of depression. The authors reported that 78.4%

(116/148) remitted during the acute phase of treatment (8–14 weeks). During the continuation phase, 15.5% (18/116) experienced relapse of major depression; thus, a total of 66.2% patients recovered fully^{47,48}. Consequently, the authors concluded that older patients with recurrent major depression can successfully be treated with a combination of antidepressant medication and interpersonal psychotherapy, and that older patients respond as well, albeit more slowly, than middle-aged patients⁴⁹.

COGNITIVE-BEHAVIORAL THERAPY

The cognitive model of depression⁵⁰ is based on the notion that, as a consequence of early learning, depressed individuals develop stable cognitive schemas or core beliefs which predispose them to negative interpretations of life events (i.e. cognitive distortions). This distorted style of thinking is hypothesized to result in depressive behavior and experience.

Cognitive-behavioral interventions for depression typically involve three active components. First is a behavioral activation component, in which the patient is exhorted to increase activities that are reinforcing, and thus increase the amount of pleasurable experience in life. Second, automatic dysfunctional thoughts are identified, explored, challenged and replaced with more accurate cognition, based on a thorough assessment of the patient's contextual environment. Third, underlying cognitive schemas or structures, which are hypothesized to drive automatic cognitive distortions and limit access to experiences that may alter these schemas, are identified and altered to more accurately reflect the patient's actual environmental, social and personal experience. Recent component analysis research suggests that behavioral activation and automatic thought modification are equally effective, and both components together are no more effective in preventing relapse than when used alone^{51,52}.

More purely behavioral interventions are derived from classic learning theory, in which problem behaviors are viewed as the result of specific antecedent stimuli and consequential events that reinforce, punish or maintain behavioral responses⁵³. This therapeutic approach views depression as a state in which there is a relative shift toward an increase in certain aversive affective reactions (respondent processes) and a concomitant reduction in the frequency of overt activities (operant extinction or punishment). In addition, histories of pervasive inescapable punishment, reinforcement of distressed behavior, classically conditioned dysphoric responses, and the evocative salience of certain stimuli depending on mood, may be examined as part of a functional analysis of depressed behavior^{53,54}. For example, a previously active person suffering from a serious illness may experience a reduction in the frequency of self-esteem-generating activities and positive social contacts, as well as increased dependency on others for the provision of positive reinforcers, and may feel bored, helpless and pessimistic. Restoration of the predictability and availability of positive reinforcers, and reduction in negative reinforcers (i.e. avoidance behaviors), is seen as linked to the curative process, with goals of symptom reduction and increased skill at identifying and obtaining appropriate reinforcement. Techniques used might include monitoring behavior and affect patterns, assigning pleasant events, stimulus control, limiting worry and depressive ruminations with time limits, behavioral exposure, and skills training (relaxation, problem-solving, interpersonal skills).

A related therapy for elderly depression, which utilizes elements associated with both cognitive and behavioral interventions described above, examines problems associated with social problem solving. Social problem-solving therapy (PST) is based

on a model in which ineffective coping under stress is hypothesized to lead to a breakdown of problem-solving abilities and subsequent depression^{55,56}. Therapeutic approaches involve identifying and modifying maladaptive beliefs or attitudes associated with ineffective problem-solving while increasing motivation to generate alternative solutions, make decisions, implement solutions and assess solution utility.

Outcome studies have supported the use of cognitive and behavioral psychotherapy in treatment of depression in elderly samples (see reviews^{1,7,33,57}). In a study comparing cognitive, behavioral and brief psychodynamic therapy to wait-list controls, Thompson *et al.*³³ found that all of the treatment modalities led to comparable and clinically significant reductions of depression. All three treatment regimens included individual treatment twice weekly for 4 weeks and weekly thereafter, totaling 16–20 sessions. Overall, 52% of the sample attained complete remission after treatment, and 18% showed significant improvement, with some enduring depressive symptoms. These rates are comparable to treatment outcomes in younger adult populations and response to pharmacotherapy^{33,58}. Follow-up research indicated that at 12 months after treatment 58% of the sample was depression-free, and at 24 months 70% of the sample was not depressed. As in acute treatment, there were no differences between treatment modalities at follow-up³⁴, although in previous research with a smaller sample size depressed geriatric patients in cognitive and behavioral therapies maintained the gains longer than those treated in brief psychodynamic therapy⁵⁹. Areal *et al.*⁶⁰ examined the efficacy of PST in a randomized controlled trial of 74 clinically depressed older adults (age 55 and over). Patients were assigned to one of three treatment conditions: problem-solving therapy (PST); reminiscence therapy (RT); or a waiting-list control. Following 12 weekly sessions, both therapies showed significant reductions in depressive symptoms at post-treatment and at a 3 month follow-up, relative to controls. However, PST showed a significantly greater number of patients, compared to RT, who were classified as improved or in remission following treatment.

Subsequent research on the same sample used in the Thompson³³ study examined the role of change expectancies relative to outcome⁶¹. Of those who were assigned to cognitive therapy, subjects who originally indicated that they expected a change from cognitive and behavioral processes attained greater improvement. In addition, some elderly patients, more familiar with the “doctor takes care of patient” mentality pervasive in medicine, may not be accustomed to the hard work involved on the part of the patients for the success of cognitive-behavioral therapy. Thompson *et al.*⁶² recommend addressing such views directly, while helping the patient to gain insight into how his/her thoughts influence mood and how new skills can help in coping with stressful events and automatic thoughts.

GROUP INTERVENTIONS

Group therapy has been shown efficacious for depression in adult samples⁶³ and in geriatric samples⁶⁴. Non-specific treatment factors that may influence positive outcomes include diffusing dependence on individual therapists, as well as providing a supportive social network. In addition, with the relative dearth of coverage for mental health care among many insurance providers, and the relative lack of parity for mental health care in Medicare, the lower cost of group therapy may be a more appealing option in older patient populations.

Beutler and colleagues⁶⁴ tested the relative and combined effectiveness of alprazolam and group cognitive therapy in a sample of 56 depressed older adults. Subjects were assigned to one

of four groups: alprazolam and weekly management sessions; placebo and weekly management sessions; cognitive therapy plus alprazolam; and management and cognitive therapy plus placebo. The cognitive therapy groups were held in 12 weekly 90 min sessions. Patients in group therapy showed a significant and consistent decline in BDI scores, while those not in group therapy failed to produce significant changes. Also, a significantly higher proportion of those in group cognitive therapy were asymptomatic at the end of follow-up (29% vs. 12%). Although the results indicate that the cognitive therapy group was more effective in reducing depressive symptoms than alprazolam, this drug (a type of benzodiazepine) is not a recommended medication for depression. However, it appears that benzodiazepines remain a relatively commonly prescribed medication in medical practice. In a sample of depressed inpatients at a large university medical center during 1993–1996, 25% of patients diagnosed with depression by a geriatric psychiatrist were prescribed only benzodiazepines by their physicians¹³. Koenig *et al.*¹³ also found that newer and older antidepressants were prescribed with the same frequency, suggesting that the prescribing practices of many physicians do not include the newer, safer antidepressants. Despite the fact that benzodiazepines are still used in clinical practice for depression, a revised research project using one of the newer antidepressants in Beutler’s design would be informative for today’s clinicians.

Steuer *et al.*⁶⁵ investigated cognitive-behavioral therapy (CBT) and psychodynamic group therapy in a sample of 33 depressed elders. The investigators assigned members to each condition on the basis of time entering the study and did not include a control group in their study design. Both treatment groups evidenced significant clinical improvement over 9 months of therapy. Of the 13 subjects who dropped out of treatment before the end of the 9-month period, 10 showed improvements in depression on the HAM-D, with mean improvement being 34%. There were no differences between treatments on the clinician-rated Hamilton Rating Scale (HAM-D), although the CBT group had lower scores on the self-report Beck Depression Inventory (BDI).

Lynch *et al.*⁶⁶ have reported unpublished pilot work using dialectical behavior therapy (DBT) skills training to treat elderly depression. Twenty-seven participants were randomly assigned to DBT skills training plus medication *or* medication alone plus clinical management. All participants were on antidepressant medication and all met criteria for MDD at baseline. DBT included weekly 2 h group skills training and weekly half-hour telephone check-in calls by a therapist for 28 weeks. Approximately 29% of the sample met strict criteria for a SCID-II diagnosis of personality disorder. Although there was a trend for DBT patients to show lower desires to please others from pre- to post-treatment relative to medication alone, there were no significant differences between treatments on outcome measures. Both DBT and medication alone showed significant reductions in HAM-D and BDI scores. Within-group analyses revealed that the medication alone group showed significant improvement over time on *only one* variable that the DBT condition did not, namely fear of sadness. The DBT condition showed: significant decreases in hopelessness and in total adaptive, avoidant, detached, and emotional coping; significantly lower sociotropy/dependency; lower desires to please others; and lower autonomy scores from pre- to post-treatment.

MAINTENANCE THERAPY AND RELAPSE PREVENTION

Despite the effective treatments available for depression in the elderly, it has been established that elderly patients who recover

from an episode of major depression are at high risk for relapse. Reynolds *et al.*⁸ used a combination of medication and psychotherapy for the acute treatment of depression in order to empirically test the most effective maintenance therapy for remitted elderly. In their study, the 107 patients who fully recovered in open acute treatment with nortriptyline and interpersonal therapy were randomly assigned to one of four maintenance therapies: nortriptyline and IPT; nortriptyline and medication clinic; placebo and IPT; or placebo and medication clinic. Combined psychotherapy (IPT plus medication) was superior (80% effect) to medication alone (57% effect), psychotherapy alone (43% effect) or placebo (10% effect) for the maintenance of treatment gains and prevention of relapse. Reynolds *et al.*⁸ concluded that combined psychotherapy and medication treatment appears to be the optimal long-term strategy in preserving depression remission and recommended that all older patients with recurrent depression be referred for psychotherapy. Research is currently under way testing the hypothesis that combination treatment is the most cost-effective way to treat recurrent depression in the elderly.

TREATMENT-RESISTANT POPULATIONS

While Reynolds and colleagues have established that combination therapy is the preferred means for maintenance of remission, there is an alarming number of elderly depressed patients who do not respond to medication, psychotherapy or the combination. Although acute remission rates of 50–70% are impressive, there remains another 30–50% of patients who do not respond. It has been found that elderly patients with co-morbid personality disorder, irrespective of level of depression, are less likely to benefit from short-term therapy than patients without co-morbidity⁶⁷ (see review⁶⁸); hence, part of the population not responding to treatment may have personality disorders. However, with the exception of case studies, no outcome study has specifically focused on treating late-life personality disorders, and those studies reporting outcomes for personality-disordered elderly have suffered from varied methodological problems⁶⁸. Nevertheless, poorer outcome and increased likelihood of relapse among personality-disordered elders, as well as continued observations that depression in the elderly is often a recurring phenomenon, require that revisions to existing treatments be made and implemented.

DYSTHYMIA

Despite the growing interest in studying treatment of depression in the elderly, we did not find any research specifically investigating the treatment of dysthymia in elderly patients. Although the research on treatment of recurrent old-age depression⁸ may reflect issues associated with treating dysthymic individuals, research protocols up to now have focused primarily on a diagnosis of major depression. However, minor depression or less severe forms of depression remain important areas of investigation. Partial responses to treatment are associated with higher rates of relapse⁶⁹ and minor depression is more prevalent than MDD and subsequently has a greater number of disability days associated with it⁷⁰. In addition, researchers have found that patients with endogenous depression (an older term with a symptom profile similar to dysthymia) responded less favorably to psychotherapy than non-endogenous depression patients, and that improvement occurred more quickly for the non-endogenous subtype⁵⁷. Thus, more research is needed to examine the public health challenges posed by old-age chronic recurring

depressive experience, partial responses to treatment, and minor depression.

CONCLUSIONS

There is a growing body of research demonstrating that psychotherapy offers significant promise for the treatment of elderly depression, preventing depressive relapse, and at times may be the preferred treatment modality in terms of both efficacy and patient choice. Referral to psychotherapy remains a problem, and current research on underdiagnosis and under-treatment of depression in older adults focuses more on describing the problem than on understanding it. However, with more insight into the reasons *why* elderly depression is inadequately diagnosed and treated, health providers can begin to develop operative means to ensure that this disabling yet treatable disorder is not ignored. General practitioners and mental health professionals do have the means to treat depression in the elderly with medication and/or psychotherapy; the work now lies in getting out into the “real world” and putting these techniques into practice.

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Long-term Management of Affective Disorders

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Depressive illness is a public health problem. One in five individuals will suffer from one form of depression or another, four-fifths of those will suffer further episodes of illness and a fifth will die by suicide or cardiovascular disorders. Affective disorders are the most commonly diagnosed psychiatric conditions in the elderly in both general and hospital practice. Their incidence and prevalence increases with age, particularly in their more severe forms, with endogenous and psychotic features. It would be true to say that non-bipolar depressive illnesses with endogenous and psychotic features are essentially illnesses of old age. Depressive illnesses are often recurrent and are associated with high continuing morbidity and mortality. Their acute and long-term management is often successful and rewarding, with the availability of highly specific and effective physical and psychological treatments.

The focus of this chapter is on the strategies of long-term management of affective disorders, starting with a review of studies of their natural history and the usefulness of antidepressants and lithium as continuation and prophylactic therapies. Particular emphasis is laid on lithium therapy, its efficacy and safety, optimal dosage and dosage regimen and important aspects of its routine in psychiatric practice.

In essence, the usefulness of any treatment with established effectiveness also applies to the elderly with depressive illness, and the evidence marshalled to support the use of these treatments often derives from studies done on younger patients. In this chapter, reference will be made to whether the study was done on younger or older patients.

NATURAL HISTORY

Studies of the natural history of depressive illness in old age have been relatively few compared with studies in younger patients. The literature on both hospital and community-based studies have been critically reviewed¹.

Post² studied patients over 60 years of age who were followed up for a period of 3 years after discharge from hospital: a quarter of the patients had lasting recovery, whilst the other three-quarters were equally subdivided into a group who had recurrences and a group who continued to be ill over the entire study period. Factors that predicted poor outcome were age over 70 and a duration of illness of more than 2 years prior to admission. Blessed and Wilson³ reported a 2 year follow-up study of a hospital cohort: one in five of their patients with affective psychosis had either not been discharged or had been re-admitted within the follow-up period. In a similar study, however, Christie⁴, found that only 6% of the patients were still in hospital within the

2 years of follow-up. The study by Murphy⁵ involving patients over the age of 65, treated in a variety of settings and followed up 1 year later, found that 43% had recovered by 1 year and 57% either had an early relapse of illness or had not recovered. Predictors of poor outcome were increased severity of illness, the presence of physical illness, longer duration of illness and the occurrence of life events during the follow-up year. Ninety per cent of patients with delusional depression had poor outcome. The study by Baldwin and Jolley⁶, using a similar methodology to that by Murphy⁵, reported a more favourable outcome: 50% of patients were fully recovered, 15% had a relapse following recovery and 18% were continuously ill. Poor prognosis was predicted by being male with physical illness, but was not associated with delusions. Longer follow-up studies over 20 years, involving elderly patients with depressive illness, showed that three-quarters of patients had recurrences^{7,8}. Remarkably, the study by Ciompi⁸ found that only 11% were totally free from psychiatric symptoms and one-third of patients had chronic depressive illness.

The year 3 follow-up of 1070 persons aged 65 and over living in the community in Liverpool⁹ showed that 72% of patients with depressive psychosis and 62% of patients with depressive neurosis were either dead or had some kind of psychiatric illness 3 years later.

In conclusion, these studies, despite differences in methodology and duration of follow-up, clearly indicate the high continuing morbidity of depression in old age: the majority of patients with depression occurring or recurring in old age will suffer further episodes of illness, with one-fifth to one-third of patients experiencing chronic illness, particularly those who present with more severe illness and those with physical conditions leading to high mortality by suicide and cardiovascular conditions (see Chapters 83, 87 on suicide and prognosis, this volume).

ACUTE MANAGEMENT

Acute management of depression in old age is often successful and rewarding, with the introduction of highly effective and specific treatment. In his review of the efficacy of antidepressants in elderly patients, Peet¹⁰ concluded that efficacy is established for drugs such as imipramine, amitriptyline, nortriptyline, phenelzine and mianserin with placebo control. Of the new antidepressants, trazodone and bupropion also show convincing efficacy in comparison with placebo. About two-thirds of patients respond well to these antidepressants. The effectiveness of psychological treatment in the acute management of depression in the elderly has not been satisfactorily established (see Chapter 79). However, reference is made to studies done on younger patients. A recent

study from the National Institute of Mental Health (NIMH)¹¹ established the efficacy of brief interpersonal psychotherapy in comparison with standard treatment with imipramine plus clinical management, particularly with patients who are more severely depressed and functionally impaired. Cognitive-behavioural therapy (CBT) was found to be less effective than these two treatments when compared with placebo plus clinical management. This study was the first comparative study of the effectiveness of interpersonal psychotherapy and CBT. Its results, however, are disappointing to those who advocate the supremacy of CBT for the treatment of depression in general and hospital practice. However, there is evidence for its long-term effectiveness in preventing relapses or recurrences of illness¹². The evidence for the efficacy of antidepressants (26 RCTs) and psychological treatment (RCTs) in older adults with depression has been critically reviewed¹³. Compared to placebo, efficacy has been established for fluoxetine, trazodone and phenelzine but no evidence for the superiority of any antidepressant over any other. In depressive patients with physical illness (18 RCTs), antidepressants were more effective than placebo. For psychological treatments in 40 controlled trials, treatments (cognitive therapy or CBT) were shown to be more effective in mild to moderate depression. However, these treatments were no more effective than non-specific attention.

The evidence for the efficacy of electroconvulsive therapy (ECT) in the severe forms of depression is overwhelming, although this derives mainly from studies done on younger patients. Of note was that the only predictor of good response to real vs. simulated (sham) ECT is the presence of delusions. There is, however, a high rate of relapse following recovery by ECT, indicating the need for continuation treatment following recovery. Post², in his 3 year follow-up study of depressed elderly patients, noted that 75% of patients relapsed when dosages of antidepressant were lowered after 3 months, and that it had been possible to discontinue treatment permanently in only 18% of patients; 38% of patients required tricyclic antidepressants (TCAs) intermittently or continuously and others had to be given lithium or had a further course of ECT. The study by Murphy⁵ had no patients on prophylactic lithium.

Continuation Therapy Following ECT

The efficacy of ECT in the management of depression in old age is often compromised by the high relapse rate following recovery. Abou-Saleh and Coppen¹⁴ critically reviewed the evidence for the efficacy of continuation therapy with antidepressants and lithium following recovery with ECT. It was shown that in four prospective placebo-controlled studies over a period of 6 months to 1 year, these drugs substantially reduced the relapse-recurrence rates during follow-up. The conclusion was that lithium would be particularly effective in those with delusional depression, bipolar depression and in the elderly.

LONG-TERM MANAGEMENT

Long-term management occurs in two phases: a continuation therapy phase to prevent early relapse of illness and a maintenance-prophylactic phase to prevent recurrence (Figure 80.1). Studies of the value of continuation therapy after recovery have established the efficacy of antidepressants and lithium in preventing relapse of illness. Continuation therapy with drugs appears to reduce the relapse rate by half compared to placebo within 4-6 months from recovery from the acute illness¹³.

The American Psychiatric Association Guidelines¹⁵ recommend antidepressant continuation therapy for 16-20 weeks following

remission, using the dose used in the acute phase. There is also evidence to support the use of specific psychotherapy during the continuation phase. The patient's clinical condition, as well as the specific treatment being provided, determine the frequency of visits. The decision to discontinue treatment should be based on the factors considered in the decision to initiate maintenance treatment, including the probability of recurrence, frequency and severity of past episodes, the persistence of dysthymic symptoms after recovery, the presence of co-morbid disorders and patient preference, in addition to consideration of the benefits and adverse effects of maintenance treatment.

In a number of controlled investigations of variable stringency, TCAs and lithium were shown to reduce the long-term morbidity and mortality in patients with unipolar illness, including the elderly. The results of these studies have, however, been disappointing, with only 48% success rate (absence of a relapse-recurrence) in the NIMH study with imipramine maintenance therapy over 2 years. The Medical Research Council study showed a success rate for amitriptyline of 32% over a period of 3 years' maintenance therapy¹⁶.

Prophylactic Treatment

Early studies indicated a poor outcome of late-life depression⁵, with high relapse, recurrence and chronicity. This view was based on naturalistic observation without monitoring the compliance, adequate dosage and duration of treatment, including prophylactic treatment, and was therefore challenged¹⁷. Controlled studies of maintenance antidepressant medication, however, showed better outcome for late-life depression¹⁸. The Pittsburgh group reported a 3 year follow-up study of maintenance treatment with nortriptyline or placebo, with or without interpersonal psychotherapy (IPT)¹⁹ and showed that 80% of patients assigned to nortriptyline, with or without IPT, remained in remission. The Pittsburgh group also identified the elderly patients who remained well after placebo-controlled discontinuation of antidepressant medication for a period of 1 year. Recovery of good subjective sleep quality by early continuation treatment was useful in identifying which remitted elderly depressed patients remained well with monthly IPT after discontinuation of antidepressant medication²⁰. Moreover, effective maintenance treatment with nortriptyline was associated with enhancement in the rate of delta-wave production in the first non-rapid eye movement (REM) sleep and of REM activity throughout the night²¹.

A recent study of the effects of treatment on the 2 year course of late-life depression²² showed a 74% survival rate without relapse. This good outcome was obtained by the use of full-dose antidepressant medication, frequent follow-up and rigorous treatment of relapses. In a prospective 1 year uncontrolled study in elderly depressed and dysthymic patients, reboxetine was shown to be effective and well tolerated²³.

The US National Institute of Mental Health consensus statement update²⁴ concluded that recent evidence supports the recommendation for at least 6 months of treatment beyond recovery for those with first onset in late life, and for at least 12 months for those with a recurrent illness²⁵. Moreover, prophylactic treatment should be of the same type and of same dosage as that which was successful in the initial acute phase. The consensus statement also concluded that treatment response and long-term outcome for all patients is generally similar to that observed in younger adults, but the temporal course may somewhat be slower in the elderly and risk of relapse somewhat greater²⁴. The use of lithium in continuation and prophylactic treatment of depression in the elderly has also been recommended^{18,26}, with the use of lower doses/plasma levels. Lithium may be a more favourable prophylactic treatment than antidepressants in recurrent

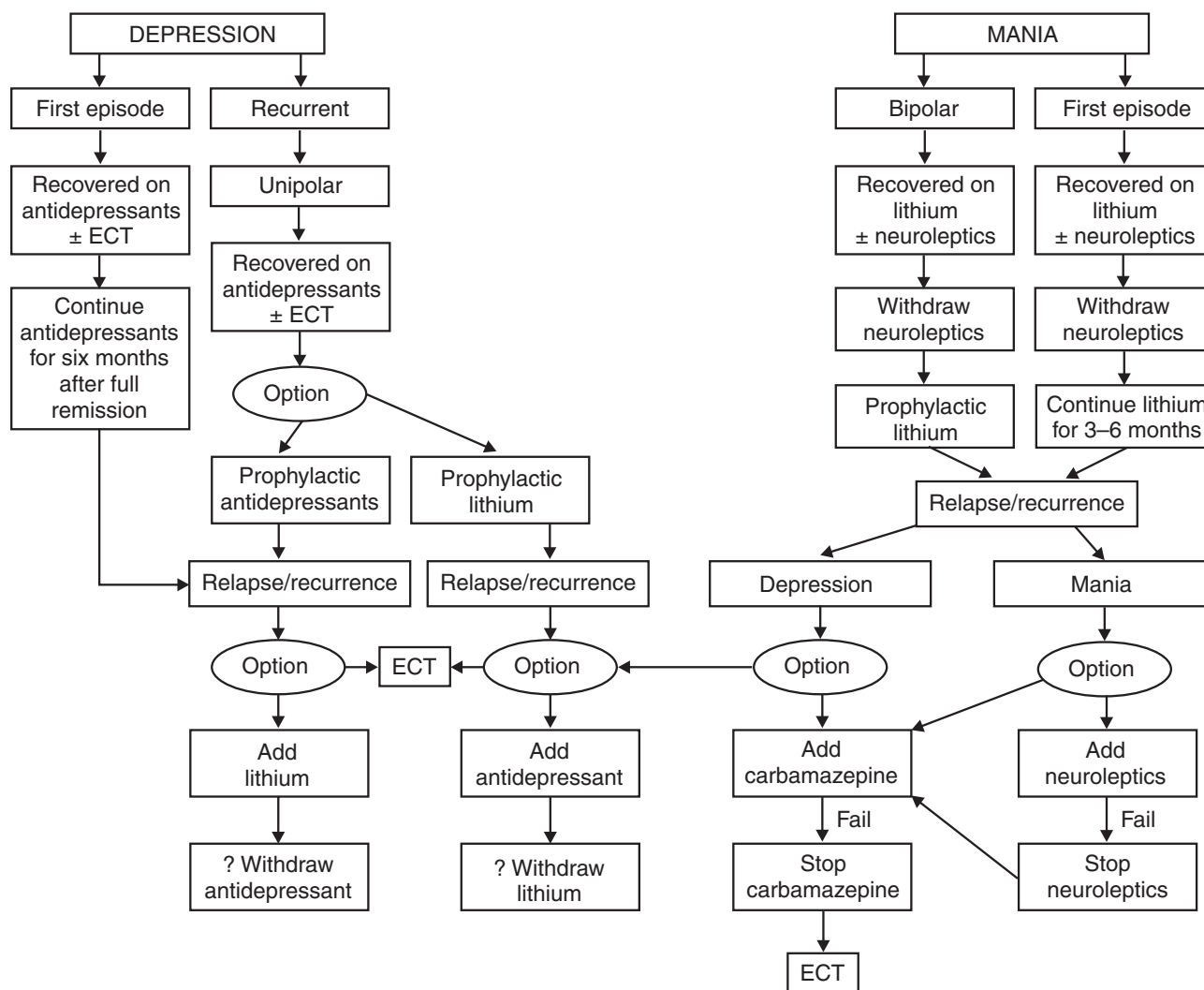


Figure 80.1 Long-term management of depression and mania

depression with melancholia and in depression with psychotic features (delusional depression), which are particularly common among the elderly, with a tendency to respond less well to antidepressants¹⁷. The other advantage of lithium therapy is the evident decreased mortality, whether from suicide or other causes²⁷. A recent 1 year prospective, placebo-controlled study of maintenance lithium in conjunction with CBT in elderly depressed patients²⁸ showed that, although CBT reduced depression severity during follow-up, lithium therapy was no better than placebo. This appears to be related to poor compliance, a finding that highlights the serious difficulties in undertaking prophylactic studies in elderly depressed patients.

Lithium Therapy

The use of lithium in the management of affective disorders has proved to be one of the most rewarding therapeutic strategies in medical practice. In the management of bipolar disorders, it has provided one of the most specific psychotropic drugs in

psychiatry. Lithium is the treatment of choice for mania and has comparable efficacy to neuroleptics, except in disturbed and agitated manic patients, who respond more dramatically to neuroleptic medication. The advantage of lithium, however, is that patients do not complain of the “strait-jacket” effect they experience with neuroleptics. Moreover, intermittent exposure to neuroleptics has been shown to be associated with a high incidence of tardive dyskinesia, particularly in the elderly, who are often less tolerant of neuroleptics and more commonly develop parkinsonian side effects. Lithium is also strongly recommended for the treatment of bipolar depression, rather than tricyclics, which may provoke hypomanic episodes and increase the risk for the development of rapid-cycling illness. There is evidence that lithium alone is effective in depression that has failed to respond to tricyclic medication, and in combination with other antidepressants in the management of resistant depression (see below).

Affective disorders are recurrent illnesses and the discovery of the prophylactic effects of lithium opened a new era in their management. In a number of controlled investigations of varying

stringency, done mostly on younger patients, lithium was shown to substantially reduce the long-term morbidity of both unipolar and bipolar disorder^{14,29}. Prospective controlled studies showed that lithium was superior to placebo and tricyclics in the prophylaxis of bipolar illness. The majority of studies evaluating its prophylactic efficacy in unipolar illness found it superior to placebo, as effective as imipramine or amitriptyline, and more effective than mianserin and maprotiline. This may be related to the greater heterogeneity of unipolar illness.

Treatment of Bipolar Disorder

Elderly patients with late presentation or late-onset mania respond well to standard antimanic treatment with neuroleptics, lithium and anticonvulsants^{30,31}. Neuroleptic treatment is best avoided in the elderly because of its known extrapyramidal side effects, except for floridly psychotic, agitated and behaviourally disturbed patients, who need rapid control of symptoms. Lithium remains the treatment of choice, followed by valproic acid³⁰. The evidence for the efficacy of lithium in late-life mania is based on retrospective and uncontrolled studies, and there have been no controlled studies of the efficacy of anticonvulsants in late-life mania. It has been suggested that valproate is a safer alternative treatment to lithium than carbamazepine, whether used as single or adjunct treatment in elderly manic patients^{30,31}. There are no guidelines regarding the optimal plasma concentration of valproate in relation to efficacy³⁰.

A recent evidence-based review of the treatment of mania, mixed state and rapid-cycling illness in younger populations³² concluded that lithium and divalproex sodium are effective in mania, whereas divalproex sodium and carbamazepine are more effective in mixed states. Divalproex sodium is the drug of choice for rapid-cycling disorder. With bipolar depression, lithium is recommended as a first-line treatment and the addition of a second mood stabilizer or a TCA would be an appropriate next step³³.

The guidelines for the continuation and prophylactic treatment of bipolar illness in late life are similar to those advocated for younger patients, except for the notion of high recurrence rates necessitating prophylactic treatment even after a first onset manic episode. Three sets of guidelines for the treatment of patients with bipolar disorder were reviewed³⁵: the American Psychiatric Association Practice Guideline¹⁵, the Expert Consensus Guideline Series (1996) and the Clinical Practice Guidelines for Bipolar Disorder from the Department of Veterans Affairs³⁶. Lithium remains the medication of choice for prophylaxis. A comparative audit of the prevalence of lithium therapy and the quality of monitoring in over-65s in Cambridge and Southampton showed a wide variation and indicated that a dedicated monitoring service leads to a better quality of treatment supervision³⁷.

Who Responds to Prophylactic Lithium?

Response to lithium varies between complete, with no further episodes of illness, to partial, with the frequency and severity of episodes reduced, to failure to respond, when morbidity continues unabated. Overall, 50–70% of patients with bipolar and severe unipolar illness show favourable responses to lithium, with a small minority who are total non-responders. The latter are often patients with rapid-cycling bipolar illness (those who suffer four episodes of illness per annum or more). The main reasons for failure of prophylaxis are poor compliance and side effects such as weight gain, increased thirst, difficulties with memory, poor concentration and loss of enthusiasm.

A recent study examined the clinical and psychological characteristics of elderly patients receiving prophylactic lithium in relation to long-term outcome of treatment (Abou-Saleh, unpublished). Elderly patients with bipolar illness had better outcome than those with unipolar illness. In their personality characteristics, those who had an excellent response showed higher scores on extraversion and energy output than those who responded less well. The most powerful predictor of long-term response, however, was their response during the first 6 months of therapy, confirming the results obtained in younger patients³⁴.

Treatment Compliance

Compliance with treatment is a major problem in the elderly. A study by Johnson³⁸ showed that the dropout rate from a group of depressed patients treated in general practice increased from 16% at the end of the first week to 68% at the end of 1 month. This was related to doubts about the benefits and less related to the occurrence of side effects. Of particular importance are the cognitive and sensory impairments of the elderly, low motivation, and poor communication by doctors of the benefits and risks to patients and their relatives. In the background lurks a nihilistic attitude and doubts about the effectiveness of treatment in both patients and doctors.

Prophylactic Lithium in the Elderly

Himmelhoch³⁹, in an open study of the efficacy of lithium in elderly patients, reported a favourable response rate in two-thirds of these patients. The majority of poor responders, however, had neurological conditions, which probably impaired the efficacy of lithium. The present author reported the results of a series of open and controlled studies of the efficacy of lithium in conventional and lower doses in the prophylactic management of affective disorders in old age⁴⁰. In the Lithium Clinic at the Medical Research Council Neuropsychiatry Laboratory in Epsom, UK, 44 male and 104 female patients with affective disorders were followed up for a period of 1–14.5 years (mean 4.9 years). They all received a slow-release lithium preparation at bedtime. Plasma lithium concentrations were maintained at 0.8–1.2 mmol/l 12 h after dosing. Prophylactic lithium was started in 47 of these patients after 60 years of age. There was no significant difference in the Affective Morbidity Index (a composite index of severity and duration of affective episodes) between the younger and these elderly patients, as shown in Table 80.1. Side effects in the older group were similar to these in younger patients. In a further study³⁴, 22 elderly patients over 60 years of age who started lithium in late life received 25–50% reduction in lithium dosage in a double-blind situation. The elderly group of patients fared as well on lower doses of lithium as younger patients and had a significant reduction in subjective side effects, such as tremor and thirst (Table 80.2). However, results for the whole group, with a majority of younger patients, showed that a reduction of 50% in daily dosage was safe; patients who had the reduced dose of lithium and plasma lithium levels of 0.45–0.59 mmol/l showed reduced morbidity during the year of follow-up, compared with the year preceding the trial, and showed fewer subjective side effects and adverse effects on thyroid and renal function.

Side Effects

Subjective Side Effects

The occurrence of subjective side effects during lithium therapy is well documented⁴¹. Side effects in the early stage (within 6 weeks

Table 80.1 General details of patients and relationship between age and morbidity during lithium therapy (results expressed as mean \pm SEM)

Age when lithium started (years)	<i>n</i>	Episodes prior to lithium	Years on lithium	AMI ^b
> 60	47	4.9 \pm 0.6	3.8 ^a \pm 0.4	0.18 \pm 0.03
40–60	79	4.2 \pm 0.3	5.5 \pm 0.4	0.17 \pm 0.02
< 40	22	4.4 \pm 0.6	5.3 \pm 0.7	0.14 \pm 0.03

^aSignificantly lower than (40–60 years) group $p < 0.01$ and (less than 40 years) group $p < 0.05$.

^bAffective Morbidity Index.

Table 80.2 Morbidity and plasma lithium level in 22 elderly patients before and during trial period (results expressed as mean \pm SEM)

Plasma lithium level (mmol/l)	<i>n</i>	AMI	
		Before trial	During trial
> 0.8	8	0.16 \pm 0.08	0.17 \pm 0.06
0.60–0.79	6	0.40 \pm 0.17	0.36 \pm 0.12
0.45–0.59	8	0.22 \pm 0.15	0.36 \pm 0.20

of starting lithium) include nausea, loose stools, fatigue, muscle weakness, polydipsia, polyuria and hand tremor.

During maintenance, weight gain, mild memory impairment and hand tremor are common complaints and polydipsia and polyuria may persist. The rate of occurrence of these side effects and their severity is related to plasma lithium concentrations. The elderly are particularly vulnerable to side effects. Smith and Helms⁴² examined the incidence and severity of side effects in elderly patients receiving lithium in comparison with younger patients. Whilst there was no difference in the total incidence of side effects, there was a trend for more serious side effects to occur more frequently in the elderly: 33% of the elderly experienced “confusion” vs. 12% of the younger group. Worthy of note is that the patients were maintained on relatively high plasma concentrations for this age group, 0.86–1.26 mmol/l. Patients with neurological conditions (parkinsonism and facial dyskinesia) develop neurotoxicity at relatively low plasma levels of less than 0.65 mmol/l³⁹. In a recent investigation, Coppen and Abou-Saleh²⁶ reported that prevalence rates of subjective side effects in elderly patients receiving low-dose lithium (plasma lithium levels of 0.52–0.6 mmol/l) were similar to those of younger patients maintained on similar plasma levels.

Thyroid Effects

Elderly patients on prophylactic lithium are more likely to develop hypothyroidism than younger ones and women are more susceptible than men. This may be related to the increased disposition of women to develop autoimmune thyroid disease in middle and old age. Indeed, pre-existing thyroid disease is the major vulnerability factor for the development of hypothyroidism during lithium therapy. Coppen and Abou-Saleh²⁶ studied thyroid function in 125 patients receiving low-dose plasma levels of lithium. Women had significantly higher levels of thyroid stimulating hormone (TSH) than men, and all four patients with abnormally high TSH levels were unipolar women. Of the 11 patients who received replacement thyroxine, 10 were unipolar patients. Thyroid function has been related to increased affective

morbidity during receipt of prophylactic lithium and has been implicated in the development of rapid-cycling bipolar disorder.

Renal Effects

Overall, 5–10% of patients on prophylactic lithium develop tubular kidney damage complicated with glomerular pathology. There is no evidence that these changes are conducive to renal insufficiency, which rarely occurs in patients who have suffered lithium intoxication or had pre-existing renal disease. Lithium dose requirements show a decrease with age, which could be accounted for by the age-dependent fall in glomerular filtration rate and lithium clearance. With age, there is a decrease in muscle mass, which limits the value of measuring serum creatinine levels as an indicator of renal function. Age-related decrease in the volume of distribution of lithium also contributes to higher plasma levels in the elderly. The daily dosage of lithium required to achieve a given plasma level may be half the dose required for a younger patient. A number of studies have evaluated renal function in relation to the dosage regimen, comparing a once per day dosage regimen to a twice-daily regimen. Contrary to the conventional wisdom, of avoiding higher peak plasma levels associated with a single daily dosage regimen, this regimen was found to be safer for renal function than a divided daily dosage regimen: functional and structural abnormalities were more pronounced in the group of patients who received lithium in divided doses than those who received it in the single daily dosage regimen, suggesting that it may be more important to have regular periods with lower levels than it is to have lower peaks.

Optimum Plasma Levels

In an open trial of lithium in elderly patients with mania (age 65–77 years), two-thirds of patients responded at levels of 0.52–0.8 mmol/l (mean 0.58 mmol/l). Two weeks after obtaining that range, two patients developed neurotoxicity and two patients only responded when the levels were raised to 0.9 mmol/l⁴⁴. Several retrospective and prospective controlled trials have evaluated the efficacy and safety of variable lithium dosage levels in bipolar and unipolar disorder⁴⁵. Some of these studies have included elderly patients. Overall, the minimal effective lithium level of the majority of patients is 0.4–0.8 mmol/l. Bipolar patients may require higher dosages/levels than unipolar patients⁴⁶. It is evident that lower dosages/levels are associated with less severe subjective side effects and adverse effects on thyroid and renal function. A prospective open study examined the ongoing morbidity of 128 patients with unipolar, bipolar and schizoaffective disorders maintained at low doses of lithium (mean level 0.56 mmol/l) over a period of 1 year²⁶. Affective morbidity was measured for three age brackets: <60 years; 66–70; and >70 years. Elderly patients aged 70+ had remarkably less morbidity than the two younger groups, but similar side effects (Table 80.3).

Shulman *et al.*⁴⁸ followed up 43 elderly patients (mean age 74 years) maintained on 12-hourly lithium levels of 0.5 mmol/l for an average period of 2 years. The majority of patients responded well to lithium, which was well tolerated, and compliance was excellent. The most common side effects were hand tremor in one-third of the patients and polyuria or polydipsia in one-quarter.

Lithium Interactions

Special consideration should be given to lithium interactions with drugs commonly prescribed for the elderly. Lithium may interact

Table 8.3 Affective morbidity and side effects in patients on lithium, divided according to age (results shown as mean \pm SEM)

Age (years)	n	AMI	BDI ^a	Side effects
< 59.9	61	0.16 \pm 0.02	6.3 \pm 0.9	8.8 \pm 1.0
60.0–69.9	51	0.14 \pm 0.02	5.8 \pm 0.7	8.7 \pm 1.3
> 70.0	16	0.06 \pm 0.01 ^a	6.9 \pm 1.7	7.6 \pm 1.4

^aBeck Depression Inventory.

with β -blockers, resulting in the slowing of heart rate, and similar interactions have been noted when lithium is combined with digoxin, with reports of increased risk of sudden cardiovascular death on this combination in predisposed patients. The elderly, however, are commonly prescribed thiazide diuretics for hypertension, which reduce lithium clearance and increase plasma lithium levels by reducing plasma volume, causing an increase in lithium and sodium reabsorption. This interaction is not observed with potassium-sparing diuretics and loop diuretics, such as frusemide. Amiloride, however, has been used for treating lithium-induced polyuria and diabetes insipidus. Finally, the widely used non-steroidal anti-inflammatory drugs have been reported to reduce lithium clearance and increase plasma levels, except for aspirin and sulindac. Ibuprofen was shown to cause significant decrease in renal lithium clearance and was linked to cases of lithium intoxication. Indomethacin, however, has been used to treat lithium-induced polyuria.

CONCLUSION

Good management calls for a comprehensive reassessment of the patient's condition to review diagnosis and to identify the reasons for treatment failure by a diligent assessment of the adequacy of previous treatments.

With regard to therapeutic strategies, these involve two concepts: alternative and adjunct therapy (Figure 80.1). The adjunct approach takes primacy over the alternative treatment approach. Adjunct treatments include lithium^{49,50}, folate and T4/T3, and also cognitive or interpersonal psychotherapy. Alternative basic treatment involves changing the conventional antidepressant to a new one, such as a selective serotonin reuptake inhibitor (SSRI) or the use of ECT. In mania, adjunct or alternative treatments are essentially anticonvulsants, principally valproate added to lithium or neuroleptics.

GUIDELINES FOR LONG-TERM THERAPY IN THE ELDERLY

The long-term management of recurrent affective disorders in the elderly starts with a careful assessment of the patient's psychiatric, physical and social condition. This involves full psychiatric examination and physical investigation for careful diagnosis, including the pattern of symptoms, previous episodes and their nature and treatment. Patients who have recovered from an acute episode of illness should be maintained for a minimum of 6 months at the same dosage, and those with a delusional first episode in late life, or who had recurrent (minimum of two episodes in 5 years) illnesses, should be considered for prophylaxis with antidepressants or lithium. The choice of antidepressant will depend on the physical condition of the patient. Conventional antidepressants should only be considered for the physically well. Otherwise, the new generation of antidepressants, including the SSRIs, should be considered. Lofepamine, fluoxetine, fluvoxamine, paroxetine and sertraline are safer antidepressants,

with no cardiotoxic effects. Fluoxetine and sertraline have been successfully evaluated as maintenance and prophylactic treatments. It is prudent to start with low doses and build up the dose gradually, with careful monitoring of side effects. Lithium is particularly effective in bipolar illness and in delusional unipolar illness, either given alone or as an adjunct in those who are already receiving antidepressants with incomplete response. Carbamazepine and valproate is a useful adjunct in bipolar patients who have failed to respond to lithium. ECT is a highly effective treatment for relapses–recurrences, whilst continuing on maintenance or prophylactic medication. The elderly require regular follow-up, to monitor their physical, psychiatric and social conditions and to deal with any emergent problems and complications, with careful attention to their social network.

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Laboratory Diagnosis: Dexamethasone Suppression Test

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There is little doubt that the application of the dexamethasone suppression test (DST) for the management of depressive illness has provided one of the most dramatic developments in biological psychiatry. Extensive investigations of this test began in the late 1960s and culminated in its introduction as a highly specific diagnostic test for endogenous depression¹. There have been hundreds of studies on its use in diagnosis, in management and as a paradigm for investigating the pathophysiology of depressive illness.

Carroll² proposed that the DST was a highly specific diagnostic test for endogenous depression, with a specificity of 96% and a sensitivity of 50%. This claim has not been substantiated by other investigators, who reported high rates of non-suppression in a variety of psychiatric conditions, including non-endogenous depression³, dementia⁴, schizophrenia and alcoholism⁵. Abnormal DST results have also been obtained in mania⁶ and eating disorders⁷. These findings have been repeatedly confirmed by other investigators and the clinical utility of the DST in the diagnosis and management of depression has been critically reviewed¹.

METHODOLOGICAL CONSIDERATIONS

Most studies have used the overnight DST, with 1 mg dexamethasone administered before midnight for cortisol estimation at 4 p.m. the following day. A cortisol of 50 ng/ml and above is taken as the criterion for non-suppression. A number of methodological aspects have been investigated, including sampling time, dexamethasone dose, the cortisol criterion for non-suppression, the assay technique and the bio-availability of dexamethasone. The criterion for non-suppression has been empirically determined by trading off the test's sensitivity and specificity, aiming for a specificity of over 90% and a sensitivity of 50%. Carroll's criterion of 50 ng/ml for non-suppression has been universally adopted, in spite of the great variation in sensitivity and specificity achieved in various centres.

Perhaps the most important contributing variable to the DST results is the dose of dexamethasone administered: the 1 mg DST has higher sensitivity and lower specificity than 2 mg DST. A corollary of the dose of dexamethasone is the corresponding plasma concentration achieved and its relationship to the results of the dexamethasone test: levels of dexamethasone in non-suppressors are lower than in suppressors⁸. The extent to which differences in bio-availability of oral dexamethasone account for differences in sensitivity and specificity of the DST for depression

is variable. The application of dexamethasone "windows" could reduce this source of test variance⁸. Dexamethasone metabolism is influenced by liver function in males and by body mass in females⁹. The combination of DST with corticotropin releasing factor has been shown to be more closely associated with hypothalamic-pituitary-adrenal (HPA) activity than DST in depressed and normal subjects¹⁰.

The influence of non-specific factors on the results of the DST has been reviewed¹; factors studied included the degree of stress, ageing, the presence of physical illness, marked weight loss, and the effects of administration and withdrawal of psychotropic drugs.

An association between increasing age and non-suppression has been shown in depressive and demented patients and in normal controls¹¹. However, a longitudinal study in elderly normal subjects followed up for a period of 2.5 years showed no effect of age and suggested that genetic factors may influence the set point of the HPA axis⁹. Physical illness is associated with non-suppression, particularly diabetes mellitus and hypothalamic disorders.

The list of drugs that interfere with the DST has been increasing, and includes hormonal preparations, liver enzyme-inducing drugs such as barbiturates, anticonvulsants and alcohol. Test results are not affected by normal doses of benzodiazepines, antidepressants and neuroleptics. High doses of benzodiazepines have been associated with normal suppression. The effects of withdrawing psychotropic medication have also been studied, with higher rates of non-suppression being observed in patients who discontinued antidepressants, neuroleptics and benzodiazepines. These non-specific factors have important effects on test results and must be ascertained when the DST is carried out in the clinical situation. They have undoubtedly contributed to the considerable variation in the rate of non-suppression in normal controls and psychiatric patients.

DIAGNOSTIC VALUE

The claim that the DST is a highly specific diagnostic test for endogenous depression (melancholia) has not been substantiated. Rates of non-suppression in endogenous depression varied (18–81%), a variation that may be partly related to the criteria used to define endogenous depression. An early review found that in 12 out of 20 studies, significant increases in non-suppression rates were found in endogenous compared with non-endogenous depressives, with the Newcastle Scale providing

better discrimination than studies using the Research Diagnostic Criteria and DSM-III criteria¹².

The Research Diagnostic Criteria for distinction between endogenous and non-endogenous depression have been validated by the DST¹³. Mitchell¹⁴ studied the DST in relation to the Core system (objective signs of psychomotor disturbance), Newcastle scale and DSM-III-R melancholia: all three definitions were associated with DST results. The Core index, but not the Newcastle scale, was associated with cortisol levels and dexamethasone levels after partialling out the effects of age and baseline cortisol levels. A meta-analysis of studies of the DST in relation to psychotic/non-psychotic depression (14 studies) and melancholic/non-melancholic depression (19 studies) reported a strong association of DST non-suppression and psychotic depression (64.1%) vs. non-psychotic depression (41%)¹⁵. The DST has good discriminating power between patients with severe psychiatric disorders, including major affective disorders, acute psychoses and dementia and patients with chronic conditions, such as dysthymic disorders (neurotic depression), chronic schizophrenia, anxiety, panic disorders and acute grief. The specificity of the test is higher in normal controls (93%) and lowest in manic patients (51%). It has satisfactory sensitivity and specificity for diagnosing secondary depression in patients with stroke. DST non-suppression after 3 months from acute stroke predicts the occurrence of depression 3 years later¹⁶. A review of nine studies reported a median sensitivity of 47% and specificity of 87%, suggesting the DST's utility for the evaluation of post-stroke depression¹⁷. However, an interesting feature is the higher rate of non-suppression in demented patients with depressive symptoms than in those without.

PROGNOSTIC VALUE

The prognostic value of the DST has been evaluated by examining the clinical outcome following antidepressive treatments in suppressors and non-suppressors, by examining changes in suppression status in relation to clinical change, and by evaluating differences in long-term outcome between non-suppressors who later converted to normal suppression and those who remained non-suppressors. Initial studies of the predictive value of the test indicated a more favourable response to antidepressant treatment in non-suppressors than in suppressors. Coppen *et al.*¹⁸, however, failed to find a difference in therapeutic outcome between suppressors and non-suppressors, classified on the basis of the traditional cut-off point (50 ng/ml) for cortisol level, but found that non-suppressors had more favourable responses to both antidepressants and electroconvulsive therapy (ECT) when the criterion for non-suppression was taken as a plasma control concentration of 100 ng/ml and above. Non-suppression confers a small advantage (1%) over suppression in predicting a more favourable response to antidepressant therapy. DST status predicted improvement on nortriptyline, but not mocllobemide or placebo¹⁹.

Changes in DST status have also been examined in relation to long-term outcome: persistent non-suppression has been associated with a greater risk of relapse within several months of treatment. Depressive patients who continued to be non-suppressors had a relapse or a fatal outcome in 77% of cases, while only 19% of those who had a conversion to normal suppression had such outcome²⁰. In depressive disorders, Coryell and Schlesser²¹ recently reported DST non-suppression to be more powerful than clinical factors in predicting completed suicides: survival analyses over 15 years showed that the estimated risk for eventual suicide was 27% in those with DST non-suppression compared to 3% in those with normal DST. In elderly depressive patients, post-dexamethasone cortisol levels

were associated with clinical improvement and high cortisolism (150 mg/ml), and predicted improvement in delusional depression²². In six out of nine studies, patients with DST non-suppression following ECT had a higher rate of relapse than suppressors²³. The value of the DST in predicting outcome following discharge has also been investigated in patients with schizophrenia and mania: non-suppressors had a more favourable clinical outcome after 6 months, and those manic patients who became suppressors before discharge had better outcomes after discharge than those who continued to be non-suppressors. In patients with senile dementia of the Alzheimer type who had depressive symptoms and who were non-suppressors, a trial of citalopram, a highly specific 5-hydroxytryptamine (5-HT)-uptake inhibitor was associated with improvement in these depressive symptoms and normalization of the DST²⁴.

COMMENT AND CONCLUSIONS

The introduction of the DST for the investigation of the psychobiology of depression has been a landmark development in biological psychiatry: it has had unprecedented evaluation and has been shown to be one of the most reproducible findings in the search for biological markers for depression. The balance of evidence indicates that the overnight 1 mg DST, with one plasma cortisol estimation in the afternoon of the next day and a cut-off point of 50 ng/ml has satisfactory sensitivity for depressive illness and high specificity when its results are compared with those obtained in normal controls and patients with minor psychiatric conditions and chronic schizophrenia. The specificity of the test for depression is unsatisfactory in comparison with other psychoses, including mania, schizophrenia and dementia. The sensitivity of the test is highest for severe depression with endogenous/psychotic features, mixed affective states and depression in old age. The test has little value in predicting response to antidepressants and ECT but provides a useful monitor of clinical progress and is a good predictor of long-term outcome: continued non-suppression or reversion from suppression to non-suppression during treatment or follow-up is an ominous sign that indicates a greater risk of relapse in the months following treatment. The influence of non-specific factors on the DST is considerable. Factors such as age, stress, weight loss, alcohol misuse, administration of anticonvulsants, withdrawal of psychotropic medication, presence of diabetes mellitus and the bio-availability of dexamethasone are important contributing factors to the test results. These factors, however, do not account for the variation in test results that appear to be essentially related to the biological process underlying depression. The 2 mg test has greater specificity, but lower sensitivity, than the 1 mg test and its diagnostic and prognostic value requires further investigation. The DST is only an aid to diagnosis and prognosis, aspects that should be essentially based on careful clinical assessment. The abnormal result provides a "physical sign" indicating biological depression. The DST test, like any other laboratory investigation, may increase the diagnostic confidence of the clinical diagnosis, but could never replace careful clinical assessment. The test is particularly useful in the assessment of biological depression associated with neurotic disorders, personality disorders, grief reaction and physical illness. Moreover, the presence of this "physical sign" indicates the necessity for vigorous physical treatment, or at least the use of a combined physical-psychological approach to such conditions as agoraphobia, obsessional illness and grief reaction, and the test may be particularly useful in monitoring the course of the illness and predicting its long-term outcome. Patients who continue to be non-suppressors are likely to relapse following recovery, and to require more intensive follow-up and continuation or maintenance of prophylactic

medication. The test may also be useful in determining when to discontinue medication: continuing non-suppression may reflect continuing vulnerability or activity of the illness, in spite of apparent recovery, and medication should only be discontinued if the test results change to normal suppression. When using the test in clinical practice, it is important to assess the presence of confounding non-specific factors and to interpret the presence of this "sign" accordingly in guiding diagnosis and prognosis. The value of the test for the clinician, like the value of a new drug, can only be empirically determined by its trial in routine clinical practice.

THE DST IN PSYCHOGERIATRIC PRACTICE

The diagnostic and prognostic value of the DST is enhanced in the psychogeriatric setting. This is related to the higher prevalence of depressive illness, endogenous type, and delusional depressions in the elderly and hence the higher prevalence of DST non-suppression (higher sensitivity) in this population. This increased sensitivity, however, is slightly impaired by the influence of age *per se* on the performance of the test, lowering its specificity; i.e. 10–20% of normal elderly patients show non-suppression. Its specificity is even lower when performed in patients with dementia, who show similar rates of non-suppression to depressive patients, which renders the DST less useful in the differential diagnosis of depressive pseudodementia. Moreover, the contribution of physical factors, including the presence of physical illness and the influence of interfering medication, further impairs its specificity for the diagnosis of endogenous depressive illness. There is, however, evidence that DST non-suppression in dementia is related to the presence of concomitant depressive symptoms^{4,25}, suggesting the usefulness of treatment with antidepressant medication. The evidence for this is, however, inconsistent: other studies have found no association between non-suppression and concomitant depressive symptoms and treatment with antidepressant medication has not consistently improved these symptoms or normalized the abnormal DST²⁶. Whether the DST has diagnostic and prognostic value in milder forms of depression, which are common in the elderly, remains to be investigated.

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Bereavement

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Advanced age and bereavement are circumstances, not psychiatric syndromes. An octogenarian may have a variety of concerns, such as remembering which medication to take when, fear of falling on icy pavement, and worry about remaining financially independent on a fixed income. A recently bereaved octogenarian is likely to have additional concerns: “Who can I share my day with?”; “Who will do the driving/cooking?”; “Where do I go from here?”. These are all essentially human rather than geriatric problems. Therefore, the psychiatry of bereavement in old age does not assume pathology or the necessity for professional intervention. Hastily prescribing psychotropic medication for a “geriatric patient” can all too easily become part of the problem, rather than the solution. Instead, we ask ourselves such questions as, “Who is this person?”; “What was the nature of this unique relationship that has been ended by death?”; “What does this loss mean to the bereaved person—and what coping skills and resources can be called upon to make it through?”.

A perspective on bereavement in old age can be developed by following the sequence of events and challenges. Fortunately, a growing number of clinical observations and research findings are available to guide this overview.

THE PRE-BEREAVEMENT SITUATION

Four questions are particularly useful in approaching an understanding of the situation that existed prior to a bereavement in old age.

1. What Was the Nature and Status of the Relationship at This Time?

This question should be answered both structurally and substantively, e.g. “He was her older brother; she relied on him for advice and support more than on any other person, including her husband”; or “She was his second wife; he had expected her to look after him—actually, to wait on him hand and foot!—and suddenly he had to be the caring person . . .”. It is useful to explore both the affective and pragmatic dimensions of the relationship. Was the survivor or the deceased heavily dependent on the other person? Had the relationship been in serious trouble? If so, was this shown by silent coexistence or stormy disputes? Did the deceased provide most of the couple’s link to society? Had the survivor organized his/her life around taking care of the deceased?

Particular sensitivity may be required in understanding an elderly person’s response to the death of an adult child or a grandchild. Recent studies suggest a highly individualized deep

interiorization of grief after the first shock of the loss^{1,2}. There may be little apparent behavioral change but much continued inner processing of the loss and an intensified life review. Researchers are reaching consensus on the multidimensionality of the older person’s response to bereavement and caution against simplistic characterizations, such as “depression”.

2. How Was the Prospect of Death Integrated into the Life Scenario?

The possibility of *anticipatory grief* should be explored, rather than assumed. Fatal accidents (especially falls and motor vehicle accidents) are far more common among people 65 and older than in any other age group (in fact, data from the US Center on Health Statistics indicate that there are more fatal accidents at 65+ than in all other age groups combined). This means that many an elderly spouse or sibling has no opportunity to prepare for the death of a person who did not seem to be in imminent danger of death. Others may have coped so long with chronic health problems that it seemed as though they might continue to go on forever. Furthermore, obvious risks may have been strongly defended against by compartmentalization or other common mechanisms. The bereavement may also take the form of the completely unexpected death of a younger family member or friend.

Despite many exceptions, however, there is a greater probability that the death will have been expected. Not only do an increasing proportion of deaths occur in old age, but modern health care has lengthened the interval between development of a life-threatening condition and the day of death. The four leading causes of death for elderly people in the USA (heart disease, malignant neoplasms, cerebrovascular disease and respiratory conditions) all involve a long period of living with progressive life-threatening conditions and, therefore, opportunity for both prolonged stress and adaptation to the prospect of death. Anticipatory grief is similar in many respects to the grief that is experienced after a death³. There are affective, cognitive and somatic components, e.g. sadness, obsession, exhaustion, etc. Clinical experience suggests that a period of anticipatory grief often helps the survivor to cope with the loss, by avoiding the element of surprise and providing the opportunity for preliminary adjustment. Traumatic grief reactions are more common when bereavement is sudden and unexpected⁴.

However, awareness of an impending loss is not an invulnerable shield against the first impact. “I thought I had cried all the tears there were to cry. I had myself under control”, one woman reported. “Not a bit of it! I went all to pieces, feeling Sammy’s hand so cold”. We are beginning to learn that anticipatory grief

has both its advantages and its risks. Did this survivor start to detach him/herself emotionally while the spouse was still alive, depriving them both of the support they needed? Or did both partners use this time to affirm their love and consult with each other in making plans and decisions?

Physicians and other caregivers have the opportunity to make valuable contributions to the survivor's ability to cope with grief by sensitive response to the situation as it exists prior to the death. Offering accurate information, suggesting other options and improving the lines of communication within the family are among the ways in which one can help to shape the anticipatory grief period into a source of strength rather than intensified anxiety. The age differential between most caregivers and the elderly bereaved people they are trying to help sometimes interferes with communication, e.g. when elders are patronized and their ability to cope with bad news is underestimated. There is also an underappreciated connection between quality of terminal care and grief recovery. A hospice physician reports that, "The pain relief we achieve for an old man in his last weeks of life helps his wife to be more of her normal self when he really needs her—and a widow with fewer regrets and nightmares later"⁵.

3. What Was the Survivor's Own Health Status Previous to the Bereavement?

This is a particularly useful question to ask with respect to the older bereaved person. The spouse was the most frequent principal family caregiver in the 40 hospices studied by Mor *et al.*⁶. About two-thirds of the spouses taking responsibility for care were people above the age of 55 and it was not unusual for the caregiver to be over 75. The elderly caregiver seldom developed new physical problems over the course of the spouse's final illness, but there was a tendency to ignore his/her existing conditions. During the first months of bereavement, the survivor's health was sometimes impaired by exacerbation of previous illnesses and impairments. It would be helpful, then, to encourage the spouse and other elderly family members and friends to look after their own health during the pre-bereavement period, and to see that health status is carefully assessed afterward. Symptoms that might appear to be part of an anxious depression syndrome could be related to physical health problems that have not received the attention they deserved.

4. Who Else Was There and What Else Was Happening?

Explored diligently, this line of inquiry may reveal significant sources of concern or potential strength that bear on adjustment to the loss. "Well, Frank's brother had come to live with us again. And he was drinking again. I could have killed him"; "The people from the church were over all the time. They really cared. We were so far from the rest of the family, but they were just like family to us . . ."; "I couldn't get anything from the doctor—what was really going on with George, what else I could do. I felt like telling him, 'I'm old—not stupid!', but you don't do that, do you?"; Some of the problems that beset the survivors after the death may be the continuation or outcome of difficulties that occurred earlier. Strained interpersonal relationships and unanswered questions create more of a burden for some survivors than the death itself.

AT THE TIME OF DEATH

Learning what happened around the time of death can help us understand the bereaved elder's state of mind. Consider, for

example, the difference between death at home and in a hospital setting. A terminally ill woman was being looked after at home by her elderly mother, with support from a local hospice service. The mother had accepted her daughter's wish to be allowed to die at home without intubation and other futile procedures. But when the daughter appeared to be actively dying, the mother panicked and called not the hospice but a visiting nursing service. Now, as a survivor, the old woman is haunted by memories of her suctioned, intubated and drugged daughter accusing her with her eyes⁷. In a more frequent scenario, an elderly person will call for assistance from paramedics when his/her spouse appears to have died. But when the paramedic team arrives, the caller may have second thoughts about seeing the spouse's body subjected to resuscitation procedures. Some bereaved elderly persons remain troubled not by the fact of the death itself, but by unanswered questions about whether or not they did the right thing at the right time.

Knowing only that the death took place in a hospital does not tell us whether supportive nurses encouraged a woman to be with her husband right through to the end and to have time with him afterward—or whether she was made to feel unwelcome and hustled away. Perhaps, again, she had not been notified until some time after the death. The particularities of the final scene can either provide an acceptable conclusion to the story of a marriage or friendship, or torment the survivor with resentment, self-doubt and other disturbing thoughts.

The psychiatrist often has the opportunity to increase the sensitivity of physicians, nurses, and other caregivers in their communication patterns around the time of death. Survivors may hold on to a word or a gesture, either as a cherished or an infuriating/depressing memory. Often one can be helpful simply by validating the survivor-to-be's feelings and giving him/her the opportunity to clarify his/her own thoughts by active listening. The generational difference between the bereaved person and the psychiatrist can be a source of misunderstanding. A widow, for example, may first respond according to the models of grief that were prevalent in her youth. It may take patience and encouragement to help her discover, express and cope with her own feelings. It is also helpful to be aware of ethnic differences in expectations for behavior around the time of death. Caregivers whose own tradition involves subdued behavior and restraint of emotions may not be prepared for families in which intense expressions of grief are expected, even required.

EARLY PHASES OF BEREAVEMENT

The idea that there are fixed "stages" of either dying or grief has attracted more believers than it deserves. One can select and force observations to fit stage theories, but to what purpose? Individual responses to grief deviate markedly from the models: this is even more common in old age, where uniqueness has been deeply engraved and polished to a high gloss.

It does make sense, though, to differentiate between responses to earlier and later periods subsequent to bereavement. Indicators of a potentially intense, disabling and protracted reaction often appear within a short time. Parke and Weiss⁸ found that those who had the most difficult time coping with the spouse's death tended to smoke and drink more heavily, use tranquilizers and express depressive mood (e.g. "Life is a strain for me . . . I wonder whether anything is worthwhile any more"). These and other investigators have found that the way the survivor responds to the loss within the first few months provides a fairly reliable forecast of what kind of adjustment will be made over a longer period of time. The obvious lesson here is that early-appearing indicators, such as loss of appetite, withdrawal from friends and activities, sleep disturbances and escape into alcohol and drugs (including

the hoarding of prescription or over-the-counter elixirs) should be taken seriously. Time alone will not necessarily prove the healer.

Dependency needs frequently come to the fore when the survivor is having great difficulty in coping. Counselors and therapists face the challenge of helping survivors to rekindle their sense of competency and autonomy. This is likely to be a step-by-step progression, in which the therapist may have to play the roles of protector and mentor for some time. The advanced age of the bereaved person does not necessarily stand in the way of therapeutic success. It has been observed in clinical practice and confirmed by research (e.g. Duran *et al.*⁹) that both the individual's personality and his/her social support often make the crucial difference in recovery from bereavement.

It is not unusual to see an elderly person appear relatively unresponsive to a death. In some situations this is the observer's failure to notice subtle but significant changes. But the lack of a strong overt response is sometimes associated with bereavement overload¹⁰. So much active and re-activated grief from previous bereavements has commanded the person's attention that there is not enough emotional energy available to respond fully to the most recent death. Survivors of the Holocaust and other disasters have often been afflicted in this way.

LATER PHASES OF BEREAVEMENT

Many senior adults have proved highly resilient after experiencing bereavement. Studies do not invariably find that widows sit around weeping and feeling helpless. The gender dimension is important here. Women outnumber men in the upper age echelons and often show themselves more skillful in providing social and emotional support. By contrast, the bereaved older man runs more risk of becoming isolated. In some environments, such as the retirement community with its high density of older women, the widower may be looked after and valued. Nevertheless, there is a major problem of hidden grief among older men, a problem that often persists for years after the loss. Quiet, keeping to himself, expressing little obvious emotion, the older bereaved man is more likely to be suffering intensely than his female peer, with her greater facility in self-expression and more sensitive support network¹¹. This not yet well appreciated fact suggests that psychiatrists and other caregivers should explore the possibility of bereavement—even a fairly remote bereavement—as the underlying factor in a variety of behavioral and somatic problems experienced by older men. Unfortunately, some of the most traumatic bereavements leave the elderly survivor with the most limited social support. This is especially the case with suicide¹². The high rate of suicide among elderly white men each year often leaves their spouses and siblings alone with their grief.

A FEW SPECIAL CONSIDERATIONS

1. Bereavement is a status marker: in spousal death, for example, the survivor is no longer a husband or a wife. From that day forward, society tends to treat him/her in a different way. We can be helpful in supporting survivors' movement to meaningful new roles and the restoration of self-confidence.
2. The "little deaths" that often accompany the later years of life can both anticipate and intensify grief reactions. The old man may have been mourning the loss of his mobility even before

his wife died; the old woman who has been relieved of her starring role as a church soloist may have felt abandoned and rejected long before her husband's death.

3. Both elders and young children are sometimes regarded as incapable of grief, although for different reasons. This unfounded assumption results in the additional suffering involved in *disenfranchized grief*¹³. Instead of being a core part of the family communication network and helping to support others in their grief, the elder may be shunted aside, thereby becoming increasingly vulnerable to depression.
3. The death of animal companions can lead to authentic grief reactions¹⁴. The grief syndrome may be very similar to what is experienced upon the death of a human companion, although the intensity is less likely to endure as long. Rage joins with grief when an elderly person is placed in a nursing home (loss of independence, loss of choice) and then learns that his/her animal companion has been taken to the pound and destroyed.

In a sense, there are no "little griefs", and there may be no "getting over" the greatest losses we experience in our lives. But most elderly adults have learned to live with disappointments, limitations and suffering. When bereavement comes, as a rule they do not need drugs, hospitalization and the whole rigmarole of the health care system. To be offered companionship and to see in that companion's eyes that one is still valued and needed is often all that is needed for the survivor to get on with life.

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Suicidal Behaviour

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This chapter concerns itself with completed and attempted suicide in the elderly. It is a major public health problem and the clinician has a crucial influence in determining outcome in suicidal patients. We shall discuss the epidemiological, social, physical and psychiatric factors involved and conclude with a consideration of preventive measures.

EPIDEMIOLOGY

Although there are significant international variations in the official completed suicide rates reported by countries throughout the world, the overall rates continue to remain among the highest in the elderly, as rates increase with age. Males aged 75 and over have the highest rates of suicide in nearly all industrialized countries, with rates for men throughout the elderly lifespan exceeding those for women. However, suicide rates in the elderly, for both sexes, have declined in recent years in many countries, with rates declining by over 30%, for example, in the UK for both sexes between 1982 and 1996¹. Explanations for such trends are largely speculative.

The prevalence of suicidal thoughts in the elderly has been investigated in some recent studies. Skoog *et al.* examined a population of non-demented 85-year-olds with the finding that among the mentally well, none had seriously considered suicide, but that the presence of mental disorder, especially major depression, was strongly correlated with suicidal feelings². The study by Forsell *et al.*³ of almost 1000 over-75-year-olds similarly found that those with frequent suicidal thoughts had a strong association with major depression. The conclusion from studies such as these is similar, notably that a careful assessment of the mental state, focused especially on the possibility of depression, is essential before any rational basis for suicidal thoughts be considered.

Little attention has been focused on non-fatal suicidal acts in the elderly, due probably to the phenomenon in recent decades being one of younger people. No countries keep national statistics, but data from centres with well-defined catchment areas allow examination of numbers, rates and trends. Cases of elderly self-harm account for about 5% of the total number of self-harm admissions to general hospitals in the UK and North America.

SOCIAL FACTORS

Studies of individual elderly suicides have drawn attention to a number of social variables. With regard to marital status,

widowed, single or divorced individuals seem to be more at risk, with marriage appearing to offer a protective factor. The great majority of elderly suicides occur in a community setting, usually in the person's home. The method of suicide varies over time, with age, gender and other sociocultural factors. It is generally found that men adopt more violent methods than women, e.g. deaths due to hanging and firearms are commoner in men. In the USA, firearms are used by over 60% of all completed suicides, with elderly White men employing this method most frequently.

The role of social isolation as a risk factor has traditionally been considered an important variable⁴, although several subsequent studies^{5,6} have found no difference between living alone and number of social contacts compared to younger suicides. Bereavement appears as a significant risk factor, with studies of attempted and completed suicide citing its relevance. The first year of widowhood seems to be a vulnerable period, with elderly widowed men being at greater relative risk.

The antecedents in terms of precipitating life events appear to differ in the elderly population compared with younger and middle-aged groups. The latter are associated more closely with interpersonal and relationship problems, financial, legal and occupational difficulties, and less with physical illness, fear of dependency and loss of function, as is often the situation in the elderly. Although complaints of "loneliness" are frequent, a recent Scandinavian study revealed that a similar proportion of younger victims (around 38%) also complained of this difficulty⁷.

PSYCHIATRIC ILLNESS

The major finding in the clinical studies of suicide and attempted suicide in the elderly is the presence of psychiatric disorder in the period prior to the event. Among the elderly, however, depressive illness is the most important predictor of suicide and this needs to be emphasized. Most comprehensive studies of completed suicide employing the psychological autopsy method report the prevalence of major depression and other mood disorders to be 60–90%. For example, Conwell *et al.*⁸, examining the relationship between age and Axis I diagnoses in a sample of 141 completed suicides aged 21–92 years, found 71% and 64% of the 75–92- and 55–77-year-old cohorts, respectively, to exhibit mood disorders, compared with 30% of the 21–34-year-old group. Major depression was diagnosed in almost 60% of the most elderly suicides, with other mood disorders accounting for 10–20% of the sample in this study. The elderly constituted the most homogeneous group, in which non-affective psychoses were rare, addictive disorders less common and late-onset depression the rule.

The symptom profile of elderly suicides prior to the event has been described in earlier studies. Barraclough⁴, in examining 30 elderly suicides, reported complaints of insomnia (90%), weight loss (75%), guilt feelings (50%) and hypochondriasis (50%) in the month prior to death. The existence of suicidal ideation often lacks spontaneous expression and it is important that such ideas are explored. Suicidal intent may be less evident, particularly where physical ill-health complaints are prominent. For example, in a recent study from Finland⁷ of over 200 elderly suicides, suicidal intent was communicated to attending healthcare professionals in only 18%. The lesson to be learnt is that the presence of somatic symptoms should not detract from a close examination of the mental state with particular regard to a depressive illness and coexistent suicidal thoughts.

Primary substance misuse disorders account for a smaller proportion of suicides than in younger age groups, with prevalence estimates of 5–40%. Similarly, non-affective psychoses are uncommonly reported compared to younger suicides. The association between suicide and dementing illnesses has received limited attention. Although advanced dementia is likely to be a protective factor, the significance of an early dementia as a risk factor for suicide is largely speculative. Individual case studies, however, indicate that in some people the fear of progressive dependency and “institutionalization” is an important dynamic, irrespective of the presence of evident cognitive deficits. There are similarly few reports on the association between personality and suicide in the elderly. Earlier studies described a personality profile of inflexibility, “failure to adjust” and poor adaptation to change. More recently, Duberstein⁹ reported a lower openness to experience (OTE) score in elderly compared to younger suicides. This profile may be summarized as a cognitive propensity to perceive problems in dichotomous, black-and-white terms, a rigidly defined self-concept and a diminished behavioural repertoire, thus decreasing the capacity to adapt to loss and change.

BIOLOGICAL FACTORS

Suicide as a distinct neurobiological entity has been investigated in the search to identify potential biological markers, although this research has been almost exclusively undertaken on a younger population. This may be partly attributable to the inherent and often contradictory data on the effect of ageing on central nervous system neurotransmitter systems. Jones *et al.*¹⁰, however, in a study of the suicidal elderly, found significant lower concentrations of cerebrospinal fluid 5-hydroxyindoleacetic acid and homovanillic acid, compared to non-suicidal and normal controls, which is in keeping with other studies in younger suicides.

PHYSICAL ILLNESS

The importance of physical illness as a major antecedent to suicide and attempted suicide in the elderly has long been emphasized. Not only does the older suicide have a higher prevalence of illness compared to his younger counterpart, but the incidence of physical illness greatly exceeds that found in the non-suicidal elderly. Several early studies reported medical illness directly contributing to suicide in around 60–70% of cases, with evidence of higher rates of physical illness among elderly males compared to females. In a recent Scandinavian study⁵, the importance of physical ill-health as a life event in the 3 months before death was demonstrated, with elderly men displaying an excess of serious somatic illness compared to elderly females (55% vs. 31%), suggesting gender differences in coping with such age-normative stressors.

Several central nervous system and systemic disorders have been linked with increased risk of suicide. These include epilepsy, multiple sclerosis, Huntington’s chorea, head injury, peptic ulcer and rheumatoid arthritis. The association of suicide with cancer is inconsistent with some studies supporting such an association, while others refute the risk, especially among hospitalized patients. In a study from Canada¹¹ involving 543 elderly suicides, with information obtained from coroners’ inquests, those with medical illnesses were significantly less likely to be referred to psychiatric services than those without a medical illness, and those with a terminal illness, comprising almost 9% of the total, were least likely of all to receive a psychiatric assessment. A number of studies have drawn attention to the importance of subjective complaints of pain prior to suicide in the elderly^{12,13}, with nearly 20% of the samples indicating it to be a major concern prior to death. The point to be reiterated is that the presence of physical illness or presentation with somatic or hypochondriacal concerns may mask the underlying depression, and this type of presentation may be of importance in elderly men, who may be less likely to verbalize their depressed mood or admit to suicidal thoughts.

These findings for completed suicide have their parallel in attempted suicide in the elderly. In a study of 100 elderly suicide attempts, 53% were considered to be suffering from significant physical illness at index assessment following the attempt¹⁴. The cohort demonstrated an increased mortality from natural causes compared to an age- and gender-matched population and, after an average of 3.5 years, 42% of the original subjects had died.

PREVENTION

Any strategy designed to prevent suicidal behaviour needs to take account of the following factors. Which individuals are likely to be at risk, how are they to be identified, and by whom? To what extent may training and education influence detection and management of vulnerable elderly individuals? How may services be improved to effect a reduction in rates of suicidal behaviour?

Risk Assessment

The act of suicide is a complex phenomenon, involving multiple psychological, physical and social factors operating at a crucial moment in the life of a vulnerable individual, and any risk assessment procedure needs to reflect these varied antecedents. A typical high-risk individual, for example, may be described as an elderly male, living alone following recent bereavement, who may have painful, chronic health problems, who is currently depressed and who has made previous suicide attempts. The problem with applying risk factors lies in the generation of high false-positive predictions associated with the relatively low base rate of completed suicide, and as yet no instruments exist with sufficient sensitivity or specificity to be clinically useful as a risk assessment scale in the elderly. It is the clinical interview that remains the cornerstone of such assessment and needs to clarify key variables. These considerations should not, however, detract the assessor from the real increased susceptibility of the elderly to eventual suicide. This can be seen particularly in elderly attempters, where suicidal intent, as measured by the Beck intent scale, is at its highest for any age group¹⁵. Attempts in the elderly are also a much stronger predictor of subsequent completed suicide, compared with attempts in younger people, with a ratio of attempts to completion estimated to be around 4:1 compared with between 8:1 and 200:1 for younger attempters. All attempts should be taken seriously.

Recognition

Despite the unique multifactorial precipitants that contribute to individual cases of suicide, opportunities for recognition exist. Suicidal intent, for example, is frequently directly expressed, albeit in different contexts, and should be taken seriously. In the recent study from Finland², for example, although nearly half of the victims, men as often as women, had brought up their suicidal ideation or intent to their next of kin, the same intent had only been communicated to healthcare professionals in 18%, despite the great majority (70%) being in contact with health services in the month before their death. The study, however, also reported that in only 24% of the cases had the healthcare professionals even asked about suicidal intentions. It is a misconception to suppose that discussion of suicidal ideas generates attempts. Most individuals feel grateful for a discussion of their suicidal feelings, about which they may feel unduly guilty.

The role of primary care services in suicide prevention is of considerable interest. Most studies report substantial levels of contact: 40–70% of elderly suicides seeing their general practitioner (GP) in the month preceding their death and between 20–50% attending in the preceding week. This raises the important issue of effective intervention at a time when the individual is particularly vulnerable. The importance of training and education programmes for GPs in the recognition and treatment of depression as a means of reducing the suicide rate arises from the Gotland Study¹⁶, in which in the year following training, suicide rates on the island fell significantly compared with other parts of Sweden, and the fall was accounted for largely by the proportion of suicides with major depression. Although not specific to the elderly, such research requires replication.

Research evidence also suggests that a minority of elderly suicides have been in contact with secondary psychiatric services prior to death. Several studies report that around 20% of their series of completed elderly suicides had contact within 6 months and around 10–15% in the preceding 1 month^{17,18}. It is important to realize that around 30–60% have no contact with health professionals prior to death, despite the high prevalence of psychiatric disorder.

The issue of treatment adequacy is of significance, given the importance of depressive illness in completed and attempted suicide, with several studies revealing inadequate or inappropriate treatment with psychotropic medication. Conwell¹⁹, for example, described the recognition and treatment of psychiatric symptoms in primary care settings for 51 elderly suicides and found only two who had received adequate treatment, with men and those with coexistent physical illness presenting the greatest challenge. Information from several Coroner's Inquest studies reveal low levels of antidepressant treatment of around 10–25%^{11,12,17}, although a more optimistic finding has been reported from Sweden, where 50% of a cohort of 75 elderly suicides had a documented history of treatment for affective disorder in the 6 months prior to death²⁰.

Preventive Strategies

Strategies for the prevention of elderly suicide have been recently reviewed from an international perspective²¹. It would be fair to say that there are limited data on the effectiveness of specific assertive outreach programmes targeting the elderly, although some encouraging initiatives have been described. DeLeo *et al.*²² described a Tele-Help/Tele-Check service for a population of 12 000 over-65-year-olds in Padua, Italy. The service provides active contacts to clients by trained staff giving information, support and prompt intervention in medical and psychological

emergencies. After 4 years of this service, only one suicide was reported, which was significantly lower than expected.

In the USA, the Gatekeepers Program of Spokane, Washington, addresses the need to contact non-self-referrals by training business personnel in the recognition and referral to health professionals of elderly distressed individuals. Such personnel include apartment managers, pharmacists, meter readers, etc. In the UK, the implementation of screening policies for the elderly, the development of community-based old age psychiatry services maintaining close links with primary care facilities, improved education and liaison links with general hospital services, and local and national audit programmes, are likely to be useful.

CONCLUSION

Although there has been a recent decline in elderly suicide rates in several Western countries, the rates remain among the highest for any age group in most societies throughout the world. Despite this, the phenomenon receives little public attention. A common assumption is that suicide in these individuals is an understandable, normal reaction to hopeless, irreversible situations and is consequently unavoidable. There is increasing debate over the individual's "right to die" and self-determination and euthanasia advocates have adopted a more prominent position.

Whatever an individual's personal views are towards the morality of suicidal behaviour, it is incumbent on the physician to pay close attention to the mental and physical state. Available evidence suggests that the great majority of individuals who attempt or commit suicide suffer from both potentially treatable psychiatric conditions (mainly depression) and associated physical and social difficulties for which much can be done. To condemn the elderly by adopting a negative approach is to succumb to the dangers of "ageism".

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Genetics and Aetiology

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The literature surrounding the genetics of psychiatric illness and bipolar disorder is expanding, but is limited and often contradictory. Great advances have been made regarding DNA analysis and mapping the human genome. Despite these advances, it is difficult to adequately analyze the available data because the genes studied are complex and exhibit significant heterogeneity in genotype and subsequent phenotype.

There is clear evidence from family studies that bipolar disorder is inherited¹⁻⁵. However, the genetic foundation of this disorder is not well understood. Leonhard's⁶ early hypothesis, that bipolar and unipolar illnesses are distinct clinical entities and do not share a common genetic vulnerability, has been supported by empirical data. However, Roglev's⁷ recent study supported the polygenic threshold model of transmission, giving credence to the hypothesis that unipolar and bipolar disorder may be one illness. Angst⁸ first demonstrated the greater prevalence of both unipolar and bipolar illness in the probands of bipolar patients, compared to unipolar probands who only showed an excess number of unipolar relatives. These data have been replicated in a number of studies⁹. Following up on earlier research demonstrating a linkage between an X chromosome marker (color blindness) and manic depressive illness¹⁰, additional chromosomal studies have linked markers on the X chromosome to bipolar disease and pedigrees from Belgium and Israel¹¹. Yet, clearly this mode of genetic transmission could not be solely responsible for bipolar illness. Families with both fathers and sons with the illness are not explained by this mode of transmission, since the father only gives the son the Y chromosome. Other researchers have demonstrated linkage between bipolar disorder and the H-ras-1 locus of chromosome 11 in a pedigree of North American Amish^{12,13}. However, separate investigations fail to confirm this linkage in Icelandic, Irish and North American¹⁴ pedigrees. Baron and Egeland later published an update and re-evaluation of their data, essentially reversing their original findings supporting linkage between chromosomes X and 11 with bipolar disorder^{15,16}. More recent studies suggest links between bipolar disorder and expanded trinucleotide repeats, potassium channel gene hKCa3, Darier's disease and velo-cardio-facial syndrome^{17,18-21}. However, other research can be found that questions some of these links²²⁻²³. Other chromosomes, including 18, 21, 22, 5, 12 and X, have been implicated in bipolar disorder¹⁷. The consensus has evolved that the constellation of symptoms making up the phenotypic expression of bipolar disorder are probably due to genetic heterogeneity, and that subtypes within the bipolar spectrum may exist.

The discrepancy in the literature can be best understood using the model of bipolar illness as a genetically heterogeneous disorder. Probably, these are complex genes with variable

penetrance that do not exhibit true Mendelian genetics¹⁷. Because of these factors, it is difficult to obtain significant "power" to demonstrate gene linkage, explaining some of the conflicting results found in the literature²⁴.

In the elderly, bipolar disorder can be divided into those patients with an early onset of their manic symptoms and those patients for whom symptoms first appear in late life, thereby underscoring possible genetic etiology vs. environmental factors contributing to the expression of manic symptoms. Mania in geriatric patients may be due to bipolar disorder, with early onset and later recurrence or chronic course, or to the first episode of mania occurring in late life. In Shulman and Post's²⁵ review of bipolar patients admitted to the hospital after age 60, the mean age of onset of the manic symptoms is 60 years. Similarly, in a group of manic hospitalized patients over 65 years old, Stone²⁶ found that 26% had no prior history of affective disorders. Other larger epidemiologic studies with patient populations of all ages have demonstrated that the onset of bipolar disorder peaks at an early age and declines throughout an individual's lifetime²⁷⁻³⁰.

Older adults presenting with the same phenotype (i.e. manic symptoms) may therefore have a very different past clinical course of their illness and may possibly fit into different subtypes of bipolar disorder (e.g. early and late onset).

Stone's²⁶ review of elderly patients with mania demonstrates that patients with a family history of affective disorder have an earlier average age of onset of their manic symptoms (53 years, compared to 60 years for patients with no family history of affective disorder). Further studies have also supported the notion that patients with early-onset mania (usually defined as less than 30 years old) have an increased number of relatives with affective disorder and possibly an increased genetic loading for bipolar disorder³¹⁻³³. However, other authors have not found a relationship between age of onset of manic symptoms and family history^{28,34}.

There is also evidence that females are more likely to develop early-onset mania^{28,35}. This evidence would provide support for the notion that genetic loading and the X chromosome may play a significant role in early-onset mania. As Winokur²⁸ points out, other studies indicate that women are more common than men in both the early- and late-onset groups³⁶⁻³⁸. However, recent epidemiological studies in the USA show approximately equal risk of bipolar disorder in males and females³⁹. Additionally, Spicer notes an increase in first episodes of mania among men but not women over 60⁴⁰.

Clinical studies would also suggest that late-onset mania is more often precipitated by organic factors than mania, which has its onset early in life⁴¹. Krauthammer and Klerman⁴² were the first to review causes of secondary mania (mania directly related to

organic illness, rather than an idiopathic cause). These researchers describe manic symptoms that were secondary to drugs, infection, neoplasia, epilepsy and metabolic disturbances. The patients in their study had a later onset of symptoms (average age of onset 41 years) and fewer family members with affective illness than is commonly reported in the literature describing bipolar subjects.

Stasiek and Zetin⁴³ have updated Krauthammer and Klerman's original study with additional organic causes of manic symptoms. These authors again emphasize the importance of looking for an organic cause of mania when there is a negative family history for affective disorder or a later onset to the bipolar disorder. Neurological disease, particularly cerebrovascular disease, is also associated with secondary late-life mania⁴⁴⁻⁴⁶. Neuroimaging of manic geriatric patients has shown differences in the basal ganglia morphology and the putamen volume when compared to control subjects^{47,48}.

McDonald *et al.* demonstrate a correlation between bipolar disorder and subcortical hyperintensities, as evidenced on magnetic resonance images in patients with the onset of manic symptoms after the age of 50 years^{49,50}. Only two of the 10 patients in this study had a family history of affective disorders⁴⁹. The significance of subcortical hyperintensities has been debated, although they are thought to result from a focal loss of brain parenchyma, due to ischemia from any cause including hypertension, vasculopathy, atherosclerosis or thromboembolism⁵¹.

Finally, Yassa *et al.*⁵² find a low incidence of organic factors precipitating mania in an elderly population. However, seven of their 10 geriatric bipolar patients demonstrated a stressful event in the 6 month period prior to their manic episode. None of these patients had a family history of affective disorders.

AFFECTIVE DISORDERS

In summary, there is evidence from family studies and chromosomal linkage studies that manic depressive illness is transmitted genetically and may have more than one mode of genetic transmission. There is also evidence from clinical studies that there may be a different etiology for bipolar disorder, which has an onset early in life, compared to those which begin in older adults. Late-onset mania is more often associated with organic precipitants and, as in other types of mania, may be closely associated with stressful events. Patients with an earlier onset to their symptoms more often have a family history of affective disorder. Clinically, patients who present with manic symptoms after the age of 50 years should be given careful consideration for an organic cause to their symptoms, particularly if there is no evidence of affective disorders in other family members.

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Epidemiology and Risk Factors

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In recent years there has been renewed research and clinical interest in the syndrome of mania in the elderly. Much of the current work in this area has been focused on the following issues: can mania first present in late life?; do the causes, features, clinical course, and responsiveness to treatment differ in older manic patients compared to younger persons with mania?; how do elderly manic patients with an early age of onset of illness differ from those with onset of illness in late life?

EPIDEMIOLOGY

The manic phase of bipolar affective disorder, or mania, is an uncommon disorder in the elderly. In the five-site Epidemiologic Catchment Center study of more than 20 000 non-institutionalized individuals, 1 month prevalence rates for mania were 0.4–0.8% for 18–44-year-olds and 0.2% in the 45–64-year-old group. Notably, no cases of mania were identified among people over age 64¹.

Nevertheless, elderly patients with mania are seen in significant numbers in a variety of clinical settings. In Roth's² retrospective review of 464 psychogeriatric patients over age 60 in a long-term hospital, 14 cases were manic. This represented 6% of the total number of cases of affective disorder. Two studies of first admissions to British psychiatric hospitals, using Department of Health statistics, found that the number of first admissions with mania either remained steady with age³ or increased with advancing age⁴. In the USA, other studies in short-stay hospitals have reported that mania accounted for approximately 5% of elderly psychiatric admissions^{5,6}. Similarly, one recent study identified 39 patients over age 60 with bipolar, manic or mixed state disorder out of 791 inpatient admissions (approximately 5%) over a 4 year period⁷. Most studies of elderly manic patients have found more females than males^{5–8} but one recent report found a slight male preponderance⁹.

For the majority of elderly bipolar patients, the first episode of affective disorder is usually a depression. Indeed, it is quite common for a first manic episode to occur 10 years or more after an initial depressive episode and to be preceded by multiple depressive episodes over many years^{5,10,11}. Generally, elderly patients have been found to have suffered more episodes of depression before a first manic episode and to have had a long gap between an initial depression and a first manic episode than young manic patients¹¹.

It must be noted that, at the present time, there is no agreed-upon standard regarding which age should serve as the dividing line between early and late onset of bipolar illness. Furthermore, age of onset itself is often difficult to determine with exactitude.

Different criteria have been used among various investigators to identify age of onset, including first onset of symptoms, first hospitalization and first time at which the patient met full criteria for the disorder⁷. Several recent studies have observed bi-modality in age of onset of mania among elderly patients. In these studies, one subgroup of patients was found to have developed bipolar disorder in early life with a mean age in their 30s, and another subgroup developed a first manic episode after age 60^{7,11}. Late-onset bipolar patients tend to have had a longer gap between first depression and first mania than early-onset bipolar patients¹¹, and in one study were more likely to be married or living with a significant other⁷.

RISK FACTORS

A number of studies have reported that elderly bipolar patients who had an early age of onset were more likely to have had first degree relatives with affective disorder than late-onset elderly bipolar patients^{11,12}. This trend holds across studies that have used ages between 20 and 60 years to divide early and late cases and suggests that genetics plays a greater role in the disease of early-onset bipolar disorder. At the same time many investigators have reported associations between late-onset mania and cerebrovascular and neurologic disease. A cohort study comparing 50 elderly patients with mania to 50 age- and sex-matched patients with unipolar depression found that 36% of the manic patients had neurological disorders compared with only 8% of the depressed patients¹³. Interestingly, among these neurologically impaired manic patients, 33% had a positive family history of affective disorder in first-degree relatives. In another recent study comparing elderly patients with early- and late-onset bipolar disorder, researchers found that patients with late-onset illness were more likely to demonstrate cerebrovascular risk factors or clinical evidence of cerebrovascular disorders⁷. In a prospective study of mania in 35 patients over age 60, the elderly manic patients had more cortical atrophy on CT scans than age-matched controls¹¹. However, no significant difference in cortical atrophy was found between elderly patients with early- and late-onset mania. In addition, subcortical hyperintensities have been reported on magnetic resonance imaging (MRI) in elderly patients with mania¹⁴. These hyperintensities are believed to be due to focal loss of brain parenchyma but they do not seem to be specific to elderly patients with mania, since subcortical hyperintensities have also been found in late-onset depression as well as late-onset paranoid disorders¹⁵.

Krauthammer and Klerman¹⁶ proposed criteria for secondary mania, which included cases of no prior family history, no prior

psychiatric history, and a definable medical or neurological etiology. While it is true that organic factors may precipitate mania in some elderly patients, these cases appear to be in the minority. Manic episodes can be caused by such widely prescribed medications as levodopa, procyclidine, pergolide, selegiline and bromocriptine¹⁷. A variety of steroids have been reported to produce manic syndromes, as have thyroid supplements, and there have been case reports of mania associated with H₂-antagonists, antiarrhythmics, estrogen and antitubercular agents¹⁸. In addition, mania can occur in association with systemic infections, such as influenza, Q fever and St Louis type A encephalitis. Cases of mania secondary to space-occupying lesions, such as meningiomas, subarachnoid hemorrhages and metastases (usually in the non-dominant hemisphere), have also been reported⁶. In these cases, mania usually resolves with removal of the offending pharmacologic agent or treatment of the underlying disorder.

More commonly, mania has been reported in elderly patients with cerebrovascular and neurological disorders. One prospective study found that 20% of patients with mania over the age of 60 had a first manic episode closely temporally related to a cerebral organic disorder¹¹. Another prospective study of 20 manic patients with onset over age 50 found that 65% developed bipolar disorder after a silent cerebral infarction¹⁹. In particular, injury to the right hemisphere appears to be strongly associated with the development of mania. There have been reports of secondary mania in patients with ischemic injury to right-sided basal ganglia, orbitofrontal cortex, and right basotemporal cortex²⁰. It has been hypothesized that these brain areas may be significant because of their connections to the limbic system and the modulation of emotion. Overall, however, mania following stroke is much less common than depression after a stroke. In one large study of 700 stroke victims only three developed manic syndromes²¹. There have been two case reports of mania secondary to infarctions in the thalamic and perithalamic areas²². Starkstein *et al.*²³ studied 11 patients who developed mania after stroke and found that eight had lesions involving limbic areas and nine had right hemispheric involvement. These patients also had significantly larger bifrontal and third ventricular brain ratios than matched control patients, indicating pre-existing anterior subcortical atrophy. Moreover, almost half of the patients had a family history of affective disorder in a first-degree relative. Taken together, the current literature on mania in patients with neurologic and cerebrovascular disorders underscores that genetic loading is also a factor contributing to the development of mania in these patients as well. Furthermore, while up to a quarter of elderly manic patients in various studies have been found to have some evidence of concurrent cerebral disease, it is still unclear to what extent these impairments play an etiological role in the development of mania. If cases in which subjects with a known previous history of affective disorder are excluded, few cases of clear secondary mania are found^{6,11,12}. An exception to this is Shulman *et al.*¹³, who felt that 36% of elderly patients they studied who were hospitalized with mania had true secondary mania associated temporally with clearly documented neurological disorders.

Mania can occur in the setting of dementia. In a chart review of 134 patients with Alzheimer's disease, 2% were found to have had mania²⁴, although others have reported higher rates²⁵. Broadhead and Jacoby¹¹ found that 32% of elderly manic patients studied scored in the demented range on cognitive testing, even though they had no history of progressive cognitive decline. Furthermore, more extensive cortical atrophy on head CT correlated with poorer test scores, but there was no significant difference between early- and late-onset manic patients with respect to CT findings or cognitive changes. It is unknown at this time whether these patients have progressed to develop true dementia. In one

retrospective study of 92 elderly patients with mania, only three went on to develop documented dementia over a 10 year follow-up period¹². To date, there does not appear to be an increased risk for elderly manic patients to develop dementia compared to the rest of the population.

As in younger patients, stressful life events have been felt by some investigators to precipitate mania in elderly patients. One study that reported on 10 elderly manic patients found that 70% had major changes in lifestyle in the 6 months preceding onset of mania. Stresses included marital discord and disruption of living arrangements⁵.

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Mania: Clinical Features and Management

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Descriptions of manic syndromes in elderly patients published during the 1960s and 1970s tended to emphasize differences in clinical features compared to younger manic patients. Post felt that elderly manic patients had more depressive features and more mood-incongruent persecutory delusions, but were less likely to have flight of ideas¹. Slater and Roth noted that euphoric states in older manic patients tended to be less “infectious” and had more expressed “hostility and resentment”².

More recent retrospective and prospective studies, on the other hand, suggest that symptoms are actually quite similar across the age span. For example, two studies have reported that hyperactivity, insomnia and thought disorder (e.g. flight of ideas) occur in 60% of elderly manic patients. Other common manic symptoms include grandiose delusions, irritability, hypersexuality and paranoid delusions^{3,4}. Similarly, a more recent study of 14 patients over age 65 hospitalized with a first manic episode found that 43% had psychotic features, including grandiose and persecutory delusions⁵. In a prospective study comparing hospitalized manic patients over age 60 to those under age 40, no differences were found between young and old patients in the time course or outcome of the manic episode. However, the authors had the clinical impression that younger patients tended to experience a more severe illness than older patients. The authors also noted that more elderly patients relapsed into a depressive episode after discharge from the hospital⁶.

There has been little research to date comparing elderly manic patients with early- and late-onset bipolar disorder in terms of clinical presentation, response to treatment or clinical course. One intriguing study of patients over age 60 hospitalized with bipolar, manic, mixed or depressed states found that elderly patients with late-onset bipolar disorder were more likely to have psychotic features compared to patients with early-onset bipolar disorder. However, early- and late-onset bipolar patients did not differ in terms of pharmacological treatment, hospital stay length, or likelihood to be admitted for a manic, mixed-state or depressed episode⁷.

Although several authors have noted that cognitive dysfunction is common in elderly patients with mania, the relationship between mania and dementia in older patients remains complex. Broadhead and Jacoby⁶ found that 32% of elderly manic patients scored within the demented range on a cognitive assessment, even though none of the patients had a history of progressive intellectual decline. Moreover, there was no significant difference between manic patients with early- and late-onset bipolar disorder in terms of cognitive impairment. In a 5–7 year follow-up study of 25 elderly patients previously hospitalized for acute mania, 32% had developed a clinically significant cognitive disorder⁸. In contrast, an earlier 10 year follow-up study of 92 elderly patients

with mania found that only 3% had developed dementia⁹. Clearly this is an area that warrants further study.

TREATMENT AND MANAGEMENT

A recent study of mental health service use by elderly patients with bipolar disorder and unipolar major depression found that the bipolar patients used more case-management services, were three times more likely to use partial hospitalization and three times more likely to have had at least one psychiatric hospitalization over the 6 months prior to the assessment¹⁰. This study highlights the importance of effective treatment and mental health monitoring for elderly patients with bipolar disorder.

As with younger patients, lithium carbonate is the mainstay of pharmacological treatment in the older manic patient. There are clear changes in the pharmacokinetics of lithium with age, due to age-related decline in creatinine clearance. In people with no renal disease there is a 30–50% decline in glomerular filtration rate (GFR) between the third and eighth decades. Since lithium is almost exclusively excreted through the kidneys, this leads to decreased clearance of lithium with age. As a result, the biological half-life in the serum increases from 18 h in adolescents to 36 h in people over age 60 and may be even longer in elderly patients with renal disease¹¹.

In addition, there is considerable agreement that both the toxic and therapeutic effects of lithium occur at lower plasma levels in the elderly compared to younger patients. Plasma levels of lithium that would be considered therapeutic for a young manic patient cause delirium in some elderly patients. The reasons behind this are unclear but it has been hypothesized that this phenomenon reflects an increase in brain sensitivity to lithium. Fine hand tremor secondary to lithium also seems to occur more frequently in people over age 60 than in younger patients. In summary, therefore, it is prudent to begin elderly manic patients on lower doses of lithium than one would for younger patients, and to aim for lower plasma levels of 0.5–0.6 mmol/l, even in the acute phase of treatment. For long-term maintenance and prophylaxis, even lower plasma levels of lithium of 0.4–0.5 mmol/l are often effective for this older age group.

Studies of patients taking lithium over many years indicate that lithium rarely causes changes in glomerular filtration rate or renal failure¹². Furthermore, lithium does not appear to decline in efficacy over the lifespan¹³. However, for a variety of reasons, some patients become unable to tolerate lithium as they age. In these cases lithium treatment must be replaced or supplemented by other mood-stabilizing medications. As with younger patients, anticonvulsants such as carbamazepine,

valproate and clonazepam are being used more frequently as an alternative or adjunct to lithium treatment¹⁴. Although systematic data regarding the efficacy and toxicity of carbamazepine treatment in elderly manic patients are lacking, there are a few reports indicating that it can be safe and effective in older patients¹⁵. The major risk of carbamazepine is the rare occurrence of aplastic anemia or agranulocytosis¹¹. However, many older patients have difficulty tolerating this medication, due to its propensity to cause sedation and ataxia.

In recent years valproate has been widely prescribed for the acute treatment of mania in both young and elderly patients with bipolar disorder. A case series study of seven older patients with long-standing bipolar disorder who had failed to respond to conventional medications reported at least minimal improvement in six out of seven patients after valproate was added adjunctively to the medication regimen¹⁶. Two retrospective studies of valproate treatment of patients over age 60 hospitalized with mania found valproate effective in improving symptoms of mania at serum levels of 31–106 µg/ml^{17,18}. In another retrospective study assessing the efficacy of lithium compared to valproate in patients over age 55 hospitalized for mania, 38% of elderly patients receiving valproate were improved at discharge compared to 67% of patients receiving lithium. However, the authors note that fewer patients on valproate therapy were within the therapeutic range and valproate serum levels of 65–90 µg/ml correlated with a better therapeutic response than did lower serum levels¹⁹.

Gabapentin is a new anticonvulsant with gaba-ergic and glutaminergic properties that has been shown to have anti-manic effects and may be effective as an adjunctive agent in the treatment of mania²⁰. It has not yet been studied systematically in elderly patients. Similarly, in a mixed-age population of patients with acute mania, olanzapine, a novel neuroleptic, was found to be more effective than placebo²¹. Novel neuroleptics such as risperidone and olanzapine have the potential to be particularly useful adjuncts in the treatment of mania in the elderly because of their lower rates of extrapyramidal side effects and tardive dyskinesia, compared with traditional neuroleptics²². As a result, they may be better tolerated by elderly manic patients than traditional neuroleptics, such as haldol, for adjunctive treatment of psychosis and agitation. More research is needed on both the short- and long-term effects of these newer mood-stabilizing and antipsychotic medications in older persons.

Finally, the clinical management of older patients with mania must always include attention to psychosocial support issues and psychotherapy. As with younger patients, individual counseling, emotional support and illness education are critical to a successful outcome. Additionally, involvement of family members and significant others in the acute treatment and long-term management of the illness helps to ensure compliance with treatment recommendations and early detection of signs of relapse.

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Prognosis

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In a follow-up study of approximately 400 bipolar patients, Grof *et al.*¹ found that virtually every patient experienced recurrence of the illness. The course of the recurring episode was predicted by the course of the previous episode. The first recurrence usually occurs 40–48 months after the initial episode, but later in the course of the illness, there may be more frequent relapses and the interval between episodes may shorten up to 50%.

Winokur², using IOWA-500 data from a long-term follow-up study, suggested that bipolar illness may burn itself out with time; thus, aging may be a mitigating factor. Cutler and Post³, however, in a review of a small number of untreated patients with severe and prolonged bipolar disorder, found a tendency for more rapid recurrences late in the history of the illness with decreasing periods of normality. In other words, if bipolar illness emerges in later years, then the episodes of mania or mania mixed with depression may tend to cluster with each other.

The course of the illness also depends on the age of onset. Mania, in general, is an illness of early onset. The mean age of onset is 30 years with a range from teens through 50 years. However, there are several patients who experience their first episode of manic illness in the later years of life^{4–7}. Unfortunately, the prognosis of mania in the elderly has not been studied as systematically as in the younger population.

Post^{8,9} suggested that the first onset of affective illness in late life carried a less favorable prognosis than early-onset patients. He further stated that elderly manic patients tend to remain disturbed somewhat longer. Although most episodes may subside in a few weeks, some may last up to 6 months. In the elderly, recurrent episodes of mania tend to occur more frequently, most commonly every 1 to 2 years, as compared to longer intervals between episodes in younger patients. Post also noted that late-onset patients displayed a significantly less frequent family history of affective disorder compared to early-onset patients.

In another retrospective study of 67 elderly bipolar patients, Shulman and Post¹⁰ found that the first manic episode occurred at about 60 years of age, often after a latency period of 10 years from the first affective symptoms. There was an average of three depressive episodes before the occurrence of the first manic episode in these patients.

More recently, Kit Stone¹¹, in a retrospective study of 92 patients admitted with mania over the age of 65, reported that 26% had no prior history of mood disorder, 30% had previously only experienced depression, half of them had at least three episodes of depression before the onset of mania and the mean latency period for the first depressive symptoms to the onset of first manic episode was 16 years. He also noted that patients with a family history of affective disorders had a significantly earlier age of onset.

In a prospective, comparative study of 35 elderly manic patients over the age of 60 and 35 young patients below the age of 40, Broadhead and Jacoby¹² reported that elderly manic patients suffered more depression before the first manic episode and had a longer interval between the first depression and the first manic episode than the young bipolar patients. They also reported clinical differences between the young and the old manic patients; although the young experienced more severe illness than the old, the elderly patients appeared to have a more fragile recovery and were more likely to relapse into a depressive episode during resolution of their manic illness. Within the elderly group, 50% of the early-onset subgroup had a family history of affective disorder in first-degree relatives as compared to only 14% in the late-onset subgroup.

In a study describing the relationship between age, signs and symptoms of mania in 40 inpatients, Young and Falk¹³ reported that increasing age is associated with attenuated manic response characterized by less intense level of over-activity and sexual drive, while thought processes are generally less disrupted. They also noted greater residual psychopathology in elderly patients as compared to younger patients over a period of time. These findings are similar to previous follow-up studies by McDonald¹⁴, Wertham¹⁵ and Lundquist¹⁶, who found that duration or chronicity of manic illness increases with age.

Ameblas¹⁷ reported a special relationship between life events and the onset of the first manic episode. This finding suggests that later in life, less stressful events may precipitate mania, and therefore late-onset manic episodes may be related to increased cerebral vulnerability in the elderly.

There has been a recent interest in the concept of secondary mania. Krauthammer and Klerman¹⁸ described secondary manias in heterogeneous groups of illnesses, including metabolic disturbance, drugs, infections, neoplasms and epilepsy. There is also increasing evidence that elderly patients with coarse brain changes secondary to stroke, head trauma and other neurological conditions appear to be more vulnerable to mania. Elderly subjects are more prone to develop these conditions and the prognosis of these secondary manias depends in part on the prognosis of the conditions causing such manic episodes.

Spicer¹⁹, in an attempt to explain the increased incidence of mania with old age (a finding not replicated in most studies), suggested that manic episodes might occur on the basis of dementia. Shulman and Post²⁰, Stone¹¹ and Broadhead and Jacoby¹², however, did not find an association between mania and dementia in elderly patients. In a retrospective clinical study of mania and old age, Shulman and Post¹² did find a temporal association between onset of mania and history of cerebral disease in 16 out of 67 (24%) of their patients. Glasser and Rabins²¹

suggested from their study that elderly men with coarse brain changes as a result of stroke, head trauma and other neurological conditions, appear to be most vulnerable to developing mania. Kit Stone¹¹ reported that 24% of his elderly patients developed mania following some sort of cerebral insult. In their comparative study of young and old manic patients, Broadhead and Jacoby¹² found that 20% of their elderly patients had first manic episodes that were closely related temporally to cerebral organic disease, in contrast to none in the younger manic group. In other words, manic episodes in late life may derive not only from a predisposition to bipolar disorder but also from cerebral pathological changes, and therefore the prognosis of the brain disease may determine the prognosis of mania in these patients.

Shukla *et al.*²² studied 20 patients with mania after head injury. They suggest a significant relationship between post-traumatic seizures and development of mania. Elderly individuals are more prone to falls and subsequently at a higher risk for developing mania after head injuries. In a recent magnetic resonance imaging (MRI) study of elderly manic patients, McDonald and Blazer²³ found an increased incidence of sub-cortical hyperintensities in the right middle third of the brain parenchyma.

Young and Falk¹³ noted a less vigorous response to lithium in older than in younger manic patients. This difference, based on age, was even more impressive since their sample maximum age was 66 with a mean of 36 years. Their data suggest that increased age may be associated with attenuation of aspects of manic psychopathology and response to pharmacotherapy. Himmeloch *et al.*²⁴, in a study of 81 bipolar patients over the age of 55, demonstrated that neurological status rather than age is the critical factor determining the natural course of the illness. Neurological status not only determined the evolution of chronic mania; it also predicted poor treatment response and lithium-induced neurotoxicity.

Despite the possible difference in the clinical presentation and response to treatments of mania in young and old age, evidence indicates a good therapeutic response to lithium for both an acute manic episode and prophylaxis in elderly patients, but close monitoring is required, with particular attention to interactions with other illness and medications. Contrary to the general belief, the prognosis of mania in old age is good and comparable to that of the young manic patient.

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The Management of Acute Mania

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Primary acute mania in the elderly (>65 years) is rare, its prevalence being much less than 1%¹. Kramer *et al.*² reported that no patients with mania were identified out of 923 elderly persons interviewed in the Epidemiologic Catchment Area study. However, other researchers³ have reported that 5%–10% of elderly patients presenting with mood disorders have mania or hypomania. Further, approximately 4%–5% of geriatric patients hospitalized for acute conditions in a state psychiatric facility were noted to have bipolar disorders⁴. Most elderly bipolar patients appear to be individuals who had an early onset of bipolar disorder and now have reached an older age. However, a first episode of mania in late life is not unknown and appears to be more commonly associated with underlying neurological/medical illnesses⁵ and the absence of family history of affective disorders⁶.

Clinicians who treat the elderly are aware that late-life mania presents unique treatment challenges. Older patients with mania who require hospitalization tend to have a slower course of improvement than younger patients⁷. This often results in longer hospitalizations and the appearance of increased treatment resistance. Further, rates of cognitive and functional impairment were found to be much higher in elderly bipolar patients over a 5 year period than seen in the general population⁸.

Complicating treatment guidelines is the observation that, as opposed to bipolar disorder in younger populations, the treatment of bipolar disorders in the elderly remains an understudied area⁹. The treatment of late-life bipolar disorder is primarily extrapolated from data obtained in studies of younger patients or mixed patient populations, uncontrolled studies of elderly patients and case reports.

TREATMENT GOALS

Management of acute mania initially begins with the treatment of the acute symptoms and assessment of cause. This can be done either in the hospital or in an outpatient setting. However, since the manic patient frequently presents with excessive energy, irritability, grandiosity, impulsiveness or even psychosis, the physician must first determine the appropriate level of care required for treatment. Usually, hospitalization is required when patients present with severe symptoms or psychosis which affects functioning and requires behavior management. This may be especially true for the patient with cognitive impairment. Hospitalization can often be useful in facilitating the evaluation of first episodes and increasing both acute and long-term medical compliance. Elderly patients in nursing care facilities may be able to be treated without hospitalization if the support system is sufficient. A hypomanic patient can usually be treated as an outpatient as long as he/she demonstrates adequate compliance and the primary caregiver does not feel overwhelmed.

Before a primary manic episode is diagnosed, a potential medical or substance-induced etiology must be ruled out. Therefore, comprehensive evaluation, physical work-up and diagnosis are necessary prior to treatment of mania in the older person. After stabilization, the focus of treatment is then directed to the maintenance phase, relapse prevention and other necessary social interventions.

PRIMARY MEDICATIONS FOR ACUTE MANIA

Lithium

Efficacy

As in younger patients, lithium is indicated in elderly manic patients for both treatment of acute manic episodes and prophylactic maintenance. Although it has been suggested that lithium may not be as effective in older patients, most studies have demonstrated that lithium is equally effective in elderly patients. However, lithium does require special precautions for use in the general adult population and especially the geriatric population. In fact, underlying dementia, cardiovascular disease or kidney disease in the elderly can increase the potential for neurotoxicity, cardiotoxicity and nephrotoxicity^{10,11}. Lithium is not considered an ideal choice for patients with these problems, despite their positive response to the drug.

Treatment Considerations

Prior to starting patients on lithium, the patient's medical history should be reviewed with special attention to systems that may be affected by lithium. Laboratory tests should include a complete blood count (CBC), serum chemistry panel [with plasma sodium, potassium, blood urea nitrogen (BUN) and creatinine levels] thyroid function tests [iodothyronine (T3), thyroxine (T4), and thyrotropin-stimulating hormone (TSH) levels] and an electrocardiogram (ECG). If the elderly patient is suspected of having renal dysfunction, a 24-hour urine collection measuring creatinine clearance is the best way of assessing glomerular filtration rate (GFR). However, if this is not feasible, a less precise estimate of GFR and renal clearance can be calculated by measurements of BUN and creatinine. Unfortunately, in patients with low muscle mass, the values may be overestimated. An electroencephalogram (EEG), computed tomography (CT) head scan or magnetic resonance image (MRI) of the brain are unnecessary unless specific neurological deficits are found in these patients on physical examination.

Dose

Older patients may be started on low doses of 150–300 mg/day, usually divided into two doses. The dose should be gradually increased according to the patient's response and ability to tolerate side effects. Lithium pharmacokinetics are affected by the aging process. As one ages, there is a decrease in total body water, renal blood flow and glomerular surface area. Therefore, the excretion of lithium decreases with age. This tendency is parallel to the reduction in creatinine clearance with age. Several studies have shown that elderly patients require one-third to one-half the lithium dose suitable for young adults¹². A longer elimination half-life (<36 h) seen in the elderly suggests that a steady state may be reached within 7–10 days of initiation of therapy. Initially, lithium levels should be monitored at least once a week, with the blood level estimated at least 12 h after the last dose (the most convenient time would be early morning prior to the first dosage).

The target for lithium concentration level should always be based on individual therapeutic response. However, studies have indicated that lithium levels of 0.8–1.2 mEq/l are generally recommended. Those studies have predominately been done on mixed-age populations. Some researchers have suggested that the elderly may respond to relatively lower serum lithium levels (0.3–0.5 mEq/l)¹³.

Side Effects

General Side Effects. Common side effects with lithium therapy are a benign fine tremor, metallic taste, polyuria, polydipsia, gastric irritability, nausea, weight gain and mild sedation. The presence of tremor is an especially common and bothersome side effect in all adults taking lithium, but it tends to increase with age¹⁴. The tremor occurs at relatively low serum concentrations, and may worsen as the concentration increases⁹. Various researchers have found the incidence of tremor in the elderly to be between 17.5–58%^{15–17}.

Lowering the dosage may reduce most of these side effects. Taking the lithium with food or using the enteric coated form can reduce the problem of gastric irritation. If lithium tremor is persistent, propranolol may be helpful, although its use should be closely monitored because of the potential for other side effects to which elderly patients may be susceptible. Low-dose thiazide diuretics may improve polyuria and polydipsia, but should be used with caution (see below).

Signs of lithium toxicity include the significant exacerbation of the above symptoms, as well as vomiting, diarrhea, ataxia, slurred speech, severe lethargy, weakness, blurred vision, severe drowsiness or, at times, agitation. Lithium toxicity is often seen in the acute treatment of a manic episode in an older adult as the lithium dose is being regulated. Therefore, frequent monitoring of blood levels and side effects is essential. If symptoms are present, lithium should be stopped immediately. Dialysis may be used to remove excessively high levels of lithium from the blood. Recovery from lithium toxicity may be prolonged in the elderly due to slow renal clearance, the presence of an underlying dementia, or possibly subclinical brain changes¹⁸.

Neurotoxicity. Excessive amounts of lithium may result in neurotoxic effects, characterized by confusion, disorientation, memory loss, ataxia and akathisia. However, several reports have shown that neurotoxic effects may develop at relatively low serum concentrations of lithium in the elderly. Himmelhoch *et al.*¹⁰ found that the presence of an underlying neurological disorder that produces parkinsonian symptoms, dementia or episodic confusion is highly predictive of lithium-induced neurotoxicity, even with low serum lithium concentrations.

Cardiovascular Toxicity. Lithium may cause both non-specific and specific ECG changes associated with repolarization. Some of the specific ECG changes are T-wave flattening or inversion (which may be caused by potassium depletion) and the appearance of U waves. Adverse effects of cardiotoxicity can present as conduction defects, irregular and slowed sinus rhythm, first degree atrioventricular (AV) block, or premature ventricular complexes (PVCS). Ventricular tachycardia is a sign of severe cardiotoxicity. Older patients are at a higher risk for these cardiovascular effects, even with milder degrees of lithium toxicity. Regular comparison of the ECG with baseline measures is important once the patient starts treatment with lithium.

Nephrogenic Diabetes Insipidus. Lithium may also cause symptoms of nephrogenic diabetes insipidus in up to 20–40% of patients on maintenance lithium therapy¹⁹. This condition is caused by the inability of the patient to concentrate his/her urine. The result is the excretion of large quantities (>3l/day) of urine, which can cause dehydration if comparable fluid intake is not maintained. This syndrome is usually reversible and is not dose-related²⁰; however, there appears to be a risk of persistent concentrating defects even after the lithium is stopped^{21,22}. Diabetes insipidus can be treated effectively with amiloride or hydrochlorothiazide (HCTZ), but each of these also has the possibility of causing electrolyte imbalance, particularly in geriatric patients.

Thyroid Abnormalities. Lithium therapy can interfere with thyroid functioning, such as synthesis, degradation and release of T3 and T4. Decrease in thyroid hormone can stimulate TSH by negative feedback. Even though compensation may occur initially, some patients develop signs of hypothyroidism²³ with goiter. If detected early, hypothyroidism is reversible, either by stopping the lithium or by the use of thyroid supplements. Lithium should be discontinued if the patient develops goiter, as this can be reversed if detected sufficiently early.

Drug Interactions

Lithium is eliminated from the body through the kidneys. Therefore any medication which may alter kidney function (especially glomerular filtration or electrolyte balance) will change lithium concentration. Medications frequently prescribed to elderly patients that may have significant drug interactions with lithium include anti-hypertensive medications [thiazide diuretics, loop diuretics, angiotensin-converting enzyme (ACE) inhibitors, and calcium antagonists], non-steroidal anti-inflammatory drugs (NSAIDs), carbamazepine, theophylline, caffeine and aciclovir. Selective serotonin-reuptake inhibitors (SSRIs) have been noted to increase the risk of lithium toxicity when used in conjunction with tricyclic antidepressants (Salama, Noveske, Vesely). Aspirin (acetylsalicylic acid) has not been shown to affect lithium concentrations.

Carbamazepine (CBZ)

Efficacy

Carbamazepine, a tricyclic anticonvulsant drug, has proven efficacy in treating adults with acute mania^{24–28}. However, all of these studies were done in mixed age adult populations. Systematic studies of carbamazepine in the elderly are lacking. Although lithium remains the preferred treatment for patients with the classic presentation of bipolar disorder, the response rate

drops to 50% when the full bipolar spectrum is considered²⁹. Greil *et al.*^{30,31} have found that carbamazepine may be useful in patients with non-classical features (mixed or rapid cycling bipolar disorder) or subgroups with depressive or schizophrenia-like features. A few investigators have identified increased benefit with carbamazepine when they treated patients who had EEG abnormalities associated with symptoms of rage and violent behavior³².

Treatment Considerations

Prior to starting patients on carbamazepine, the patient's medical history should be reviewed, with special attention to previous episodes of blood dyscrasias or liver disease. Laboratory tests should include a complete blood count (CBC), serum chemistry panel, and liver function tests [lactate dehydrogenase (LDH), ALT, AST, bilirubin and alkaline phosphatase levels].

Dose and Administration

Treatment in older patients should be initiated with low doses, starting with 200–400 mg/day by mouth. The dose should be increased gradually and slowly. Carbamazepine therapeutic levels are in the range 4–12 ng/ml; however, these parameters are based on the anticonvulsant effect. There is no standard blood level recommendation for the treatment of bipolar disorder. The ideal therapeutic blood level is subjectively determined and depends on the patient's clinical response and the appearance of side effects. During the initial 6–8 weeks of treatment, blood levels may decrease despite maintaining a regular dose, due to carbamazepine's ability to induce its own metabolism. Checking blood levels and adjusting appropriately is important for maintenance of the treatment effect. Geriatric patients may be able to maintain adequate therapeutic levels on lower doses (400–600 mg) due to slower metabolism.

Side Effects

Bone marrow depression (aplastic anemia, decreased cell counts, agranulocytosis) and hepatotoxicity can occur with carbamazepine. Risk of bone marrow depression is estimated at 1/125 000. Weekly monitoring of blood counts, liver profile and carbamazepine levels is indicated during the early weeks of treatment. Idiosyncratic reactions are rare; however, patients should be asked to report immediately reactions such as fever, sore throat or severe fatigue of sudden onset.

The most common adverse effects of carbamazepine are dizziness, drowsiness, vertigo, ataxia, diplopia, nystagmus and blurred vision⁹. These effects are usually dosage-related and may be minimized by reducing the dose. Skin rash, urticaria, Stevens–Johnson syndrome, Lyell's syndrome, photosensitivity and dermatitis have also been reported, so that carbamazepine should be discontinued if these symptoms appear.

Other side effects include cardiovascular complications, such as congestive heart failure, hypertension, hypotension, arrhythmias, AV block and thrombophlebitis³³ as may occur with other tricyclic agents. The risk is greatest if patients already have underlying cardiovascular disease.

Drug Interactions

Carbamazepine interacts with a variety of other medications. Serum carbamazepine concentrations are markedly increased

when carbamazepine is given with SSRIs, erythromycin, isoniazid, calcium channel blockers and danazol. Blood levels will also increase to a lesser degree with cimetidine, valproate, phenobarbital, phenytoin and theophylline. Carbamazepine decreases neuroleptic blood levels, which may lead to a lower efficacy of the neuroleptic in combined therapy.

Valproate (Valproic Acid)

Efficacy

Valproate is an anticonvulsant also shown to be effective in the treatment of bipolar disorder in the general adult population. It has also been documented to be effective for the elderly bipolar patient^{34–44}. Valproate primarily works through the GABA system, and has recently been used to treat patients with acute mania who are resistant to traditional drugs such as lithium and carbamazepine^{36,45}. It remains the drug of choice in patients who have failed to respond to lithium or carbamazepine or who are unable to tolerate these medications because of their side effects. Valproic acid may be used effectively in combination with lithium if there is only a partial response to either drug. Like carbamazepine, valproate causes minimal cognitive side effects, which is of particular advantage in the elderly patient.

Treatment Considerations

Valproate appears to be very well tolerated by elderly patients. Prior to starting patients on valproate, liver function tests and a CBC should be performed. Albumin levels should be performed on the frail elderly. Valproate may be used in combination with lithium, although experience suggests that a trial of monotherapy is usually indicated initially⁹.

Dose and Administration

Valproic acid is available in three forms: valproic acid, sodium valproate and divalproate sodium. Gastric irritation is not uncommon with valproic acid or sodium valproate treatment. In such patients, the use of divalproate sodium may reduce gastrointestinal (GI) irritability. This medication is available in three strengths, 125 mg, 250 mg and 500 mg. Treatment should begin with low divided doses (125 mg twice a day) and be gradually increased to 500–1000 mg/day, depending on tolerance, side effects and blood levels. Most studies evaluating the effectiveness of valproate in bipolar disorder have used the target drug concentrations found effective in the treatment of epilepsy (50–100 mg/ml)⁴⁶. Higher blood levels may sometimes be required to achieve optimal clinical response.

Valproate is rapidly absorbed from the gastrointestinal system and has a half-life of 6–16 h in younger patients, although this could be longer in geriatric patients. In serum it is predominantly protein-bound after absorption. The decrease in serum albumin levels associated with aging may result in lower dosages being required for elderly patients to achieve adequate blood levels. It is metabolized in the liver prior to elimination.

Side Effects

Gastrointestinal side effects of valproate include nausea, vomiting and indigestion. Occasionally abdominal cramps, constipation and diarrhea are reported. Anorexia has also been observed, although weight gain appears more common. These effects are

usually transient, diminishing with continued treatment. Persistent GI symptoms may respond to a reduction in the dose or changing to the enteric-coated form.

Common central nervous system (CNS) effects are sedation and ataxia, which are generally dose-related. Nystagmus, headache, diplopia and asterixis are less common side effects. Dizziness and lack of coordination can occur and are also dose-related.

Minor elevations of liver transaminases (SGOT, SGPT and LDH) can occur as side effects of treatment. Hepatotoxicity with hepatic failure has been reported in children but less frequently in adults. Baseline and periodic monitoring of hepatic function is indicated during the initial phase of treatment.

Idiosyncratic reactions may cause alopecia, which can be treated effectively with vitamin supplements containing zinc and selenium. Other side effects, such as thrombocytopenia, leukocytosis, edema, weakness and skin rash, have been reported but are generally rare.

Drug Interactions

Because valproate is extensively metabolized in the liver, it may interact with other medications. Valproate may inhibit the metabolism of phenobarbital, ethosuximide, phenytoin and some tricyclic antidepressants, resulting in higher blood levels of these drugs. Frequent monitoring of blood levels and appropriate adjustment of dosage may be needed to maintain therapeutic blood levels during the initial stages of treatment. Combinations of valproate and clonazepam may produce lapse of memory, which could present a significant problem in elderly patients. The use of anticoagulants such as aspirin and warfarin should be closely monitored, as the potency of these drugs may be enhanced.

Newer Anticonvulsants

Gabapentin

Gabapentin is a novel anticonvulsant agent structurally related to γ -aminobutyric acid (GABA), currently approved as an adjunctive therapy in patients with partial seizures. Its mechanism of action is not yet fully understood. Gabapentin has been increasingly used for the treatment of bipolar disorder, behavioral disturbances in Alzheimer's disease, and social phobia. Multiple case reports and small open-label studies in the general adult population have demonstrated gabapentin to be effective as monotherapy or as an adjunct therapy for the treatment of acute mania, or as a prophylactic therapy for bipolar illness⁴⁷⁻⁵². Experience in the non-demented elderly bipolar patient is limited.

Gabapentin is not metabolized and is not protein-bound. It is excreted in the kidneys essentially unchanged. It has few pharmacokinetic interactions with other medications. Some researchers have reported a small increase in Depakote levels when used together⁵³. Gabapentin is not associated with any hematologic or hepatic problems and does not require monitoring of serum concentration. In addition, gabapentin has a relatively benign side-effect profile. The most commonly reported adverse effects are sedation, dizziness, ataxia and fatigue⁵⁴. These are usually minor and transient. These properties have made gabapentin a very attractive medication choice for use in bipolar patients who are receiving multiple medications, experience blood or liver problems, or in whom blood level monitoring is a problem⁵⁵. Some researchers⁵¹, however, have suggested that gabapentin may only exert a "moderate" antimanic effect, and that its onset of effect may be delayed when compared with other mood stabilizers. Thus, gabapentin may be less effective in the

acute treatment of mania, and is recommended as adjunctive therapy in severe mania.

The effective dose of gabapentin in the treatment of bipolar disorder is not yet known. Most trials in young adults have used a dosage range of 300-2400 mg/day, although the use of larger doses has also been documented. Ferrier⁵⁵ has suggested that gabapentin may be particularly efficacious in rapid-cycling bipolar disorder (dose range 1500-2400 mg/day). Elderly patients would presumably require less, due to age-related decreases in creatinine clearance. The usual starting dose is 300-600 mg/day in divided doses.

Lamotrigine

As the use of anticonvulsant medications for treatment of bipolar disorder has proved successful, other anticonvulsants have also been increasingly used. Lamotrigine is an anticonvulsant approved as an adjunctive treatment for refractory epilepsy. The exact mechanism of action is not yet known, although evidence suggests that it reduces the release of excitatory amino acids (by blocking voltage-dependent sodium channels) and may act as a calcium channel antagonist^{56,57}. Early studies and case reports in young adults have suggested that lamotrigine may be useful as a monotherapy or as an adjunctive therapy for bipolar disorder^{58-61,95,96,98} and is particularly effective in rapid-cycling bipolar disorder and bipolar depression⁶²⁻⁶⁶. Lamotrigine has not been studied in elderly bipolar patients.

Lamotrigine is metabolized in the liver, with a half-life of 25-30 h. Protein binding is 55%. Lamotrigine does not appear to induce the P450 system and it has little interactions with other psychotropic medications. However, carbamazepine may decrease lamotrigine levels while valproate may increase them.

The most common side effects are headache, nausea, diplopia, dizziness and ataxia. A skin rash may occur in 5% of patients, more commonly with older age, rapid escalation of dose and the concomitant use of valproate. In most cases the rash is mild and transient; however, a few patients have developed Stevens-Johnson syndrome. Therefore, the medication is usually titrated very slowly (beginning at 12.5 mg/day) and is discontinued if a rash develops. Lamotrigine has also been reported to cause confusion or psychosis or to induce mania. There are no current standard dosage recommendations and routine measurement of serum concentrations is not required.

ADJUNCTIVE MEDICATIONS FOR ACUTE MANIA

Clonazepam

Efficacy

Clonazepam is a nitrobenzodiazepine derivative indicated in the treatment of absence seizures, infantile spasms, myoclonus and atonic seizures. It has proven efficacy in reducing seizure frequency and has also been used effectively in restless leg syndrome, panic disorder and Tourette's disorder. Clonazepam has also been used with success in treating acute mania^{67,68}. It has also been used to augment mood stabilizers during the acute treatment of manic episodes, allowing a decrease in the use of neuroleptics in non-psychotic mania⁶⁹⁻⁷¹. Clonazepam has not been well studied for use in geriatric patients with bipolar disorder.

Dose and Metabolism

The antimanic effect is attained in younger adults with a dose range of 2-16 mg/day in divided doses⁶⁷. The half-life in younger

adults is 20–80 h. Older patients require much lower doses for clinical efficacy because of delayed metabolism and therefore a much longer half-life. A clinical response is usually seen in the dose range 2–6 mg in divided doses in such patients.

Side Effects

The most common side effects are ataxia, disinhibition, drowsiness and sedation. A paradoxical effect (excitation, agitation, irritability) has been observed, and is more likely to occur in patients with underlying neurological illness. Clonazepam can potentiate other sedating drugs, such as antihistamines and alcohol.

Benzodiazepine use is always recommended as short-term therapy only. Patients maintained on clonazepam may develop tolerance and dependence after 6 months of consistent use. The use of benzodiazepines in the elderly is of special concern, due to the potential to cause decrease in cognition, alertness or balance. Clonazepam should always be gradually tapered to avoid withdrawal symptoms.

Lorazepam

Efficacy

Lorazepam is a short-acting benzodiazepine that has proved effective in the treatment of acute mania^{72,73,97,99}. Lorazepam is frequently used as an adjunctive medication for the control of symptoms of acute mania until the mood-stabilizer becomes effective. Like clonazepam, the use of lorazepam has decreased the use of neuroleptics for agitation during the acute manic episode^{74,75}. This medication is not indicated for maintenance treatment. The use of lorazepam in the elderly bipolar patient is not well studied.

Dose and Metabolism

Lorazepam is available in both oral and intramuscular forms, and absorbed rapidly with either route of administration. It is the most common benzodiazepine used in geriatric populations due to its short half-life (10–20 h), large therapeutic index, no metabolites, gluconide conjugation and rapid onset of action. The dose range is generally 1–4 mg/day in divided doses for a geriatric population, titrated by individual response side effects.

Side Effects

Like all benzodiazepines, increased and persistent sedation may increase the risk of falling, especially in the elderly. Paradoxical behaviors and disinhibition can occur in a small percentage of patients, most commonly those with underlying neurological illness. Lorazepam also potentiates other sedating drugs; therefore, close monitoring is required when using lorazepam in the elderly.

ANTIPSYCHOTIC MEDICATIONS

Acute manic episodes of moderate to severe degree, or manic episodes associated with symptoms such as hallucinations, delusions, paranoia or severe irritability or agitation, can be treated initially with antipsychotic medications. Most typical antipsychotic medications effect clinical improvement by blocking

dopamine pathways in the brain and provide sedation through antihistamine effects. Atypical antipsychotic medications are thought to effect clinical change through a combination of dopamine-blocking effects and serotonergic activity. Clozapine, risperidone and olanzapine have been reported to be effective in the treatment of acute mania, both as single agents and in adjunctive treatment^{76–81}. These studies, however, have not involved elderly patients.

High-potency typical antipsychotics tend to cause significant extrapyramidal symptoms, such as rigidity, bradykinesia, tremor, dystonia, akathisia and a Parkinson-like syndrome, but a low incidence of hypotension, cardiovascular toxicity and sedation. Low-potency typical antipsychotics cause significant sedation, postural hypotension and peripheral anticholinergic effects. The newer atypical antipsychotics have less severe side effects, but still may cause sedation, orthostasis or extrapyramidal side effects. Given the high risk of such side effects in geriatric patients, low-potency neuroleptics should generally be avoided. The atypical antipsychotics are increasingly being used, due to their preferred side-effect profile. The use of antipsychotic medications in conjunction with other medications appears to have limited adverse interactions; however, a few studies have reported the development of neurotoxicity from the combined use of typical antipsychotic medications and lithium^{82–84}.

Geriatric patients typically require lower doses than middle-aged patients. Lower doses also minimize important side effects, particularly the risk of tardive dyskinesia in the long-term use of neuroleptics. Once acute symptoms improve, the antipsychotic dosage can be lowered or even discontinued while stabilizing patients on antimanic medications.

ELECTROCONVULSIVE THERAPY (ECT)

Electroconvulsive therapy (ECT) has been shown to be a highly effective treatment for acute mania^{85–87}. Several studies of both geriatric and general adult manic patients have shown an improvement in approximately 80% with ECT^{88,89}. This is especially significant, since ECT is frequently used for patients who have been resistant to other treatments or who have significant medical co-morbidities.

Contraindications

Even though there are no absolute contraindications for ECT, the risk of morbidity and mortality is increased in certain conditions. These include space-occupying intracerebral lesions or other conditions that may increase intracranial pressure, unstable vascular aneurysms or malformations, intracerebral hemorrhage, recent acute myocardial infarction or severe uncontrolled hypertension⁹⁰. However, if ECT is required, risks can usually be minimized by pharmacologic treatment during ECT.

Risks and Adverse Effects

Usually, elderly patients tolerate ECT very well. The mortality rate for elderly patients is 0.01%, roughly the same as for the anesthesia induction itself^{88,91,92}. Two-thirds of the deaths are from cardiac complications, such as ischemia, arrhythmias and transient severe increases in blood pressure. Most incidents occur immediately after the treatment or in the recovery period.

Side effects commonly observed during ECT include confusion and short-term memory loss. The latter is generally temporary, but occasionally some patients complain of prolonged loss, although no organic or irreversible changes in the brain have

been found⁹³. The risk of confusion and memory loss tends to be associated with high stimulus intensity, bilateral electrode placement, increased number and/or frequency of treatments, older age and pre-existing cognitive deficiencies. Adjustment of the stimulus waveform, decreasing the stimulus intensity, using unilateral electrode placement or increasing the interval between treatment may decrease the cognitive side effects.

Other somatic side effects include headaches, nausea, and muscle soreness. Prophylactic treatment with analgesics or using an increased dose of the muscle relaxants during ECT may minimize these symptoms. ECT patients are also at a higher risk for falls.

Treatment Considerations

Prior to initiating ECT treatment, a focused medical history and physical examination is necessary to assess and minimize any potential risk factor. Laboratory tests should include a hematocrit or hemoglobin, serum electrolytes and an electrocardiogram (ECG)⁹⁰. Most practitioners also obtain either an EEG, CT of the head, or brain MRI prior to ECT. If a patient has a history of musculoskeletal disease or osteoporosis, spinal X-rays may be obtained to evaluate the presence of underlying compression fractures, so that anesthetic/muscle relaxant medication may be properly adjusted. Dental evaluation is of particular importance in aged patients and attention should be given to patients who have loose teeth or none or only partial dentures.

An informed consent should be obtained prior to initiating ECT. In geriatric patients whose judgement and insight are compromised by their illness, the family should be involved in treatment decision making and consent. In the case of patients not competent to give consent, the legal guardian must be identified and approve consent.

The use of psychiatric medications during ECT should be minimized. Lithium should be discontinued prior to ECT, since it may increase the risk of status epilepticus and prolonged muscular blockade with succinylcholine. The anticonvulsants (valproate, carbamazepine) should also be discontinued, since they inhibit the ECT seizure. It should also be remembered that all benzodiazepines increase seizure threshold. They can be tapered prior to ECT or a benzodiazepine antagonist can be used just prior to the procedure.

Early reports suggested that mania was more resistant to ECT or required more frequent treatments than depression. Recent research has found this not to be true. The patient's clinical improvement is the best guide in deciding the optimal number of treatments. An average of 6–12 treatments is usually required for optimal response. Recommendations for treatment parameters may be found in several other texts^{90,91,94}.

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Part G

Schizophrenic Disorders and Mood-incongruent Paranoid States

Late-life Psychotic Disorders: Nosology and Classification

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The notion that mental disorders could occur in particular periods of life, such as old age, appeared only in the second half of the nineteenth century¹. During this short history, there has been little agreement, and much confusion, regarding the nosology and classification of late-life mental disorders, particularly in the area of psychotic disorders.

One group of individuals with late-life non-affective psychotic disorders consists of those with onset of a psychotic illness (e.g. schizophrenia, delusional disorder) early in life, who have now aged. Debates about the classification of these patients have followed the same lines as those regarding younger adults (e.g. degree of overlap between psychosis and affective disorders, issues of duration and course) but most would agree that these patients should retain their original diagnosis into old age unless symptoms change dramatically enough to warrant a different diagnosis. A second group of patients consists of those individuals with new onset of non-affective psychosis late in life. In a majority of such cases, psychosis is secondary to a general medical condition such as dementia. There has been little controversy about the classification of these “organic” psychoses in the elderly, and this group of individuals will not be discussed further in the present chapter. In contrast, the classification systems for late-onset non-affective, non-organic psychoses are replete with overlapping terms, multiple definitions of the same nomenclature and different methods of conceptualizing similar disorders. Only recently, with the publication of an international consensus statement², has at least the geriatric psychiatry community come to some agreement on a set of terms to describe late-onset psychotic disorders.

EARLY HISTORY

Kraepelin was one of the first clinical researchers to recognize that non-affective psychoses could arise in middle age or later in life. Although the term “dementia praecox”, with its inherent emphasis on an early age of onset, would seem to exclude late-onset cases, Kraepelin himself reported that one-third of his patients had symptom onset after age 30³. Kraepelin also studied a group of patients he described as suffering from “paraphrenia”. This term had been used earlier by Guislain as a synonym for the syndrome of “folly”⁴ and was used by Kraepelin to characterize a group of patients with minimal volitional and affective disturbance, prominent paranoia and a relatively preserved personality. While some of the subgroups of paraphrenia he described had a relatively later age of onset,

Kraepelin did not consider paraphrenia to be exclusively a late-onset disorder. Furthermore, follow-up studies of these patients showed that they did not differ greatly from those classified as dementia praecox^{5,6}. Thus, many of Kraepelin’s followers ultimately came to believe that dementia praecox and paraphrenia were the same disorder and that this disorder could arise early or late in life.

Other clinician investigators working during this time, however, felt that psychotic disorders arising for the first time in late life should be classified separately. The history of these classifications has been thoroughly reviewed⁷. Gaupp⁸ distinguished between dementia praecox and a disorder diagnosed for the first time in post-menopausal women that was characterized by depressive agitation, resulting in “mental weakness”. Stransky⁹ used the term “dementia tardiva” to describe late-onset dementia praecox. Some authors emphasized the prevalence of paranoid symptoms among those with onset of psychosis late in life by using terms such as “paranoia chronica”¹⁰ or “involutional paranoia”¹¹. Following this tradition, Albrecht’s¹² classification of late-onset psychotic patients distinguished between patients with paranoid symptoms and little personality disturbance (“presenile paraphrenia”) and those with “depressive madness resulting in imbecility”. The latter category seemed somewhat similar to a late-onset form of dementia praecox. Others who described syndromes of late-onset dementia praecox used the terms “involutional paraphrenia”¹³, “stiffening involutional psychosis”¹⁴ and “paraphrenia”¹⁵. Unfortunately, the use of “paraphrenia” to indicate an age of onset distinction led to a great deal of later confusion. Some psychiatrists employed the term to indicate a separate phenomenology independent of age of onset (much like Kraepelin’s original use; e.g. Leonhard¹⁶), while others used that diagnosis to encompass most late-onset psychoses.

1940–1970

Using his father’s term for dementia praecox, “schizophrenia”, Manfred Bleuler¹⁷ described individuals with “late-onset schizophrenia” as those with onset after age 40 exhibiting symptoms similar to those with an earlier onset of the disorder and no evidence of brain disease. Very few of these patients had onset after the age of 60. This classification was adopted by most subsequent German authors⁷.

In the UK during this period, however, the classification of late-onset psychotic disorders took a somewhat different path. Studying a group of patients with onset after age 60, Roth and

Morrissey¹⁸ described a syndrome of paranoid delusions and hallucinations in the context of preserved intellect, personality and affect. Because of the phenomenological similarity to Kraepelin's "paraphrenia" and due to its late onset, Roth and colleagues termed this disorder "late paraphrenia"^{19,20}, a name that was designed to encompass all late-onset, non-affective, non-organic psychoses in which paranoid symptoms were prominent. Thus, the term was both broader than late-onset schizophrenia, in that it encompassed late-onset delusional disorder, and more restrictive, in that it did not include non-paranoid forms of late-onset psychosis. Post²¹ developed a different descriptive system. He divided late-onset (after age 50) psychoses into paranoid hallucinosis, schizophreniform syndrome, and schizophrenic syndrome. Based on a 3 year follow-up, however, he concluded that these three diseases were actually a continuum of the same disorder with slightly different symptom profiles.

European debates and developments were slow to influence the classification system used in the USA. The first *Diagnostic and Statistical Manual of the Mental Disorders* (DSM-I)²² used the term "involuntary psychotic reaction", which encompassed both paranoid ideation and depression in older patients. This amalgam of affective and psychotic symptoms in the elderly was split in the second edition (DSM-II)²³ in favor of "involuntary paranoid state (involuntary paraphrenia)" and "involuntary melancholia". The former disorder, like Roth's late paraphrenia, was characterized by "delusion formation with onset in the involuntary period . . . The absence of conspicuous thought disorders typical of schizophrenia distinguishes it from that group"²³. Schizophrenia could be diagnosed in individuals with any age of onset.

1970-PRESENT

As European psychiatrists began to study patients with late paraphrenia more systematically, new classification systems in the USA were restricting the diagnosis of late-onset psychosis. One of the five Feighner Research Criteria²⁴ for schizophrenia was age of onset before age 40. In the third edition of the DSM (DSM-III)²⁵, a diagnosis of schizophrenia could not be made if the onset of symptoms was after age 45. Late-onset psychosis that involved persistent persecutory delusions with prominent hallucinations could be given a diagnosis of "paranoid disorder". This classification system was in stark contrast to both earlier RDC criteria²⁶ and to the 9th version of the *International Classification of Diseases* (ICD-9)²⁷, neither of which imposed age-of-onset restrictions for schizophrenia. The ICD-9 also allowed for a diagnosis of paraphrenia at any age. The revised version of DSM-III (DSM-III-R)²⁸ rectified the omission of late-onset schizophrenia by providing a separate diagnostic category for those diagnosed with schizophrenia after age 45. In the most recent version of the DSM (DSM-IV)²⁹ and the ICD (ICD-10)³⁰, no special categories exist for late-onset psychoses, although schizophrenia may be diagnosed at any age.

TOWARD A CONSENSUS

It is clear from this historical review that there has been little consensus regarding the classification of late-onset non-affective non-organic psychoses. Two opposing lines of thought have pulled the terminology in different directions. On the one hand, some authors have preferred to emphasize the similarity of late-onset psychoses to the corresponding early-onset disorders. This has resulted either in the use of terms such as "late-onset schizophrenia" and "late-onset delusional disorder" or has prompted a move toward ignoring age of onset altogether in

classification (e.g. DSM-IV, ICD-10). On the other hand, some members of the psychiatry community (mainly those in the UK) have preferred to emphasize differences between the phenomenology of late- and early-onset psychosis and thus have tended to use distinct terminology, such as "paraphrenia" or "late paraphrenia".

Thus, questions remain about which terminology would optimally serve the clinical and research communities. There are at least two overlapping issues to consider. First, how similar or different is the late-onset, non-affective, non-organic psychosis from early-onset disorders? If late-onset patients are no different from early-onset patients in terms of demography, phenomenology, etiological factors, prognosis and treatment, then it would be redundant to classify them in a separate category. If, however, such features differ between early-onset and late-onset individuals, then it would seem important to preserve a distinct diagnostic category in order to encourage further research and allow for optimal prognostic evaluation and treatment. The magnitude or extent of the differences between early- and late-onset individuals should also influence the terminology chosen for the diagnostic categorization. If a majority of critical clinical features are shared with an early-onset disorder, then it would make sense to adopt a term such as "late-onset schizophrenia". If the extent of differences is sufficiently large, a separate term would be warranted. A second issue to consider in determining the best classification scheme is what age of onset should be called "late". Most of the American studies of late-onset non-affective, non-organic psychosis have included patients with onset after 45 and generally before age 65. In addition, among the patients in Bleuler's late-onset schizophrenia studies, only 4% had an onset after age 60⁷. In contrast, most studies of late paraphrenia have been conducted with patients whose onset was after age 65. Differences in age-of-onset between late-onset schizophrenia and late paraphrenia studies may help to explain some of the diagnostic confusions that have persisted.

Only recently has the weight of evidence become sufficiently great in the field of late-life psychoses to allow for adequate consideration of these issues. In July 1998, the International Late-Onset Schizophrenia Group met to present reviews of published data on late-onset non-affective, non-organic psychosis and to develop a consensus statement regarding diagnostic categories². The statement recognizes two illness classifications: late-onset (onset after the age of 40 years) schizophrenia and a very-late-onset (onset after 60) schizophrenia-like psychosis. Thus, the group determined that it was important to recognize a diagnostic distinction based on age of onset, due to differences between late- and early-onset patients, but that the disorders were not sufficiently different to warrant a separate nomenclature. In addition, the group felt that a further distinction was warranted within late-onset patients between those with onset in middle age and those with very late onset, based on major differences between these groups.

The similarities and differences among early-onset schizophrenia, late-onset schizophrenia and very-late-onset schizophrenia-like psychosis are summarized in Table 89.1. There are many areas of similarity between both late-onset groups and early-onset schizophrenia, such as symptoms³¹⁻³³, family history³², brain imaging findings³⁴⁻³⁶, and the nature of cognitive deficits³⁵. The decision to retain the word "schizophrenia" in the nomenclature of both disorders was driven by these strong similarities. On the other hand, the consensus statement's distinction between those with middle-age-onset and old-age-onset psychoses was motivated by epidemiological, etiological and symptom differences between these two groups. Very-late-onset schizophrenia-like psychosis is different from both early- and late-onset schizophrenia, in that these cases tend to be associated with sensory impairment and social isolation²⁰, are less likely to exhibit formal thought disorder and

Table 89.1 Comparison of typical-onset (age 15–40) schizophrenia, middle-age-onset (age 41–65) schizophrenia, and very late-onset (age > 65) schizophrenia-like psychosis

	15–40	41–65	> 65
Female:male ratio	0.6:1	2:1	up to 8:1
Poor premorbid functioning	++	+	–
Family history of schizophrenia	++	++	–
Sensory deficits	–	–	+
Negative symptoms	+++	++	–
Thought disorder	+++	+++	–
Strokes, tumors	–	–	+
Neuroleptic dose	+++	++	+

+, presence; –, absence; number of symbols indicates degree of presence or absence.

affective blunting but more likely to have visual hallucinations^{26,34,37} and have less familial aggregation of schizophrenia⁴⁰. It should be emphasized that the members of the International Late-onset Schizophrenia Group were not unanimous in their support of the particular age cut-offs given in the consensus statement and also felt that the proposed nomenclature was not an end but a beginning of future research into this important topic.

Late-onset schizoaffective disorder and late-onset delusional disorder are not specifically addressed in the consensus statement. Based on recent research³⁸, late-onset schizoaffective disorder appears to share a majority of critical clinical and demographic features with late-onset schizophrenia. Thus, late-onset schizoaffective disorder appears to be a subgroup of late-onset schizophrenia in which mood symptoms are also present. Late-onset delusional disorder, by contrast, can be distinguished from late-onset schizophrenia by a unique preoccupation with non-bizarre delusions in the context of preserved affective and personality functioning in other domains²⁹. In addition, treatment of these individuals may be more challenging than in schizophrenia due to a difficulty in establishing rapport with therapists³⁹. Cognitive function, however, is somewhat more preserved in older patients with delusional disorder than in those with schizophrenia³⁹. Unfortunately, there is a lack of research comparing early- and late-onset delusional disorder.

Further research is needed to clarify the classification of late-onset psychotic disorders. Specifically, longitudinal follow-up studies are necessary to determine whether the course of illness is different in the three groups of patients and how the course of late-onset disorders compares to that of early-onset syndromes. Such an enterprise is greatly aided by the consensus classification recently proposed. In summary, despite a tumultuous history, the future for research and clinical work in late-onset psychotic disorders appears to be on firmer footing for the new millennium.

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Clinical Assessment and Differential Diagnosis

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The schizophrenias of late life, previously called late paraphrenia¹ and paranoid states in old age, present fascinating, complex, biopsychosocial problems that span the whole of psychiatry and medicine. The characteristic features of late-onset schizophrenia have been summarized by several authors²⁻¹⁰, although never better than in the seminal paper by Kay and Roth².

These conditions usually present with paranoid ideation, most commonly persecutory in nature, with or without other schizophrenia-like symptoms. The central abnormality implied by the term "paranoid" is a morbid distortion of beliefs, but not all distorted beliefs are delusions and not all elderly people expressing them are mentally ill. Many aspects of old age increase vulnerability, exposing elderly people to abuse and victimization, and a sensitive appreciation of this situation is needed when assessing the paranoid elderly patient.

Furthermore, not all patients with delusions have schizophrenia¹¹ and the first aim of assessment will be to clarify the nature of paranoid symptoms and consider a differential diagnosis. In most cases a diagnosis will be clear from the detailed historical account of symptoms and their course, abnormalities of mental state and simple physical investigations. Assessment must, then, evaluate the individual's level of functioning, independence, vulnerability, social and family support and physical health, which will all be of relevance to aetiology and management. The assessment will aim to consider patients and their symptoms within the wider context of their social environment, physical and psychological limitations. It is, therefore, necessary to have knowledge of premorbid personality, life style, life experience and cultural background, remembering that young and older generations have important cultural differences.

Ideally, the assessment will take place in the patient's home, when environment may be maximally appreciated. Home assessment provides a more complete picture of the patient's circumstances and helps put the problem into a living context. Commonly, paranoid ideas in old age relate to the patient's immediate, local environment and people within it. Herbert and Jacobson³ used the term "partition delusions" to describe the belief that things were happening just the other side of the wall, floor or ceiling. Post¹² found that paranoid symptoms often temporarily disappeared when the patient was removed from the hostile environment and this can give a misleading impression of their nature.

The floridly deluded and hallucinated patient is easily recognized, but in old age paranoid ideation may be almost plausible when complaints of being abused, victimized, stolen from or manipulated are not beyond the bounds of possibility. Trying to establish the validity of such claims requires observation and information from a variety of sources.

Table 90.1 Differential diagnosis

Delirium
Dementia
Organic delusional/hallucinatory disorder, secondary to physical illness or drugs
Late-onset schizophrenia
Delusional disorder
Depression
Mania
Schizoaffective disorder
Paranoid personality disorder
Factual (basis in fact)
Sensory impairment

We need, first, to consider the differential diagnosis of paranoid symptoms (Table 90.1) and how clinical assessment helps to differentiate diagnostic categories before discussing the process of assessment in more detail.

DIFFERENTIAL DIAGNOSIS

Christensen and Blazer¹³ found the prevalence of paranoid ideas in a community sample to be 4%. Leuchter and Spar¹⁴ retrospectively reviewed 880 psychogeriatric admissions and the 8% who suffered a first episode psychotic illness met DSM-III criteria for major affective disorder (36%), organic mental disorder (43%) and primary paranoid disorder (21%). The more common conditions will be reviewed briefly from the point of clinical differentiation, although for detailed consideration reference should be made to the relevant chapters.

Delirium (Acute Confusion)

The history is short, usually days or a few weeks, and the onset rapid. Paranoid ideas and hallucinations occur in 40–50%¹⁵. These are typically poorly organized, fluctuating and variable in content, while hallucinations most commonly occur in the visual modality. Other features of delirium will normally be present.

Dementia (Chronic Confusion)

Ballinger *et al.*¹⁶ found delusions and hallucinations in 38% and 34% of 100 dementia admissions. The study by Burns *et al.*¹⁷ of

178 Alzheimer patients revealed persecutory ideation (20%), delusions (16%) and hallucinations (17%) to be common. Fifty per cent of patients with multi-infarct dementia may have delusions at some time¹⁸ and the clinical course of diffuse Lewy body disease is particularly characterized by paranoid ideation and hallucinations¹⁹.

The manifestations of progressive, global, cognitive impairment will usually be present, although dementia may present with paranoid symptoms that can be indistinguishable from functional illness²⁰. Paranoid ideas are frequently related to cognitive deficits, especially memory, leading to accusations of theft¹⁷ or problems arising from perceptual difficulties and misidentification⁸. Like delirium, these fluctuate and may be ferociously denied, or forgotten, at interview, although the theme and content remain fairly consistent.

Depression

If depression is of delusional proportions, biological and characteristic depressive symptoms are usually marked. Delusions and hallucinations, occurring in all sensory modalities, are normally mood-congruent but incongruent symptoms occur and may be difficult to distinguish from those of primary paranoid disorders.

Kay *et al.*²¹ suggested six historical variables that help distinguish affective and paranoid psychoses: life events and family history of affective illness favoured an affective diagnosis, while low social class, few surviving children and social deafness favoured paranoid disorder. Premorbid personality proved the best discriminator, with paranoid patients being solitary, shy, touchy, suspicious and emotionally aloof, and patients with affective disorders reporting subjective ratings of high premorbid anxiety.

Mania

Traditional teaching suggested that mania in old age was both rare and atypical in presentation. Broadhead and Jacoby's²² prospective study found that young and older-onset patients were clinically very similar. The onset of mania in old age is more common than once thought²³ and the majority of patients will have a history of affective disorder, some 50% having had three or more depressive episodes, with a latency of 15–17 years from first depression to mania^{22–25}.

Organic Delusional/Hallucinatory Disorder

Paranoid hallucinatory disorders have been associated with a variety of organic conditions^{8,11,26–29} and pharmacological agents^{5,8,11,30}. The symptoms may be typical of functional disorders^{26,31} and the diagnosis depends on establishing a clear aetiological link and temporal relationships between a physical disorder or drug and mental disturbance. As Kay³² put it, "Had the organic diagnosis not been reached independently of the psychiatric symptomatology, most of the cases would have been regarded as, indubitably, schizophrenic".

The more common causes encountered in clinical practice include hypothyroidism, intra- and extracerebral tumours, epilepsy and cerebrovascular disease, and pharmacological agents such as psychostimulants, anti-parkinsonian and dopaminergic drugs and steroids. Alcohol intoxication and withdrawal from alcohol, benzodiazepines and barbiturates may all cause paranoid states, and withdrawal syndromes should be particularly considered when psychosis develops shortly after a hospital admission.

Paranoid Personality Disorder

This is necessarily a life-long problem which must be demonstrable from early adulthood. It is characterized by a sensitive and defensive attitude that causes people to feel they are victims of life and interpret events in a self-referential way¹⁴. The effects of ageing and the vicissitudes of later life may accentuate these traits and, if dementia or functional illness supervene, will colour the symptomatology.

Late-onset Schizophrenia

Kay and Roth's² description of the characteristic features of this condition has never been surpassed. Schizoid and paranoid premorbid personality, reduced likelihood of marriage and fertility, living alone with few surviving relatives, and deafness contributing to social isolation, and a limited but significant hereditary predisposition for schizophrenia with female preponderance, all characterize this disorder. The whole range of psychopathology typical of schizophrenia may be evident, although personality is more often preserved and negative features less prevalent^{2,3,20,33,34}. Roth and Kay³⁵ provide a thoughtful discussion of the apparent similarities and differences of the associated features of late- and early-onset schizophrenia.

Delusional Disorders

These are conditions characterized by a persistent, circumscribed delusional theme and if hallucinations occur they are not prominent. They are defined by their delusional content, which may be erotic, jealous, hypochondriacal, persecutory or grandiose. These conditions have not been the subject of systematic study in old age, when they are thought to be relatively rare⁸. Onset is usually in middle age but as patients normally function well outside their particular delusion and symptoms frequently persist they may present in old age. Unlike late-onset schizophrenia, delusional disorder seems not to be associated with premorbid paranoid personality or deafness³⁶, although querulent paranoia has been related to deviant personality structure⁶⁵. Familiarily they appear unrelated to affective or schizophrenic illnesses^{37,38}. Howard *et al.*³⁹ found dilatation of lateral and third ventricle volumes by magnetic resonance imaging (MRI) to be more a feature of delusional disorder than schizophrenia in old age, as defined by ICD-10 criteria.

A small retrospective study comparing paraphrenia (schizophrenia of late onset) with paranoia (delusional disorder of late onset) found cerebral infarction on CT brain scan to be a feature of paranoia rather than paraphrenia. Furthermore, social isolation and being unmarried was not a feature of paranoiacs, with cerebral infarction suggesting separate groups defined by organic or social associations. Response to antipsychotic drugs was worse for paranoia⁴⁰.

ASSESSMENT

Interview

Interviewing paranoid elderly people may be complicated by deafness, speech problems or visual handicap, so time and patience are essential. An informant history is mandatory and often several sources may be required.

It is crucial to establish the interview situation, explain its purpose, allay anxieties and put patients at their ease. The patients should decide whether they prefer to be seen in private or with a

confidant(e) as another's presence may equally inhibit or encourage the disclosure of sensitive material. For similar reasons an informant may wish to speak privately though discussions should never appear clandestine.

Deafness and communication problems should be openly acknowledged, hearing aids worn and working, and extraneous noise eliminated, otherwise false impressions of cognitive state may be formed⁴¹. If a patient is seen in a hospital setting, insist on a separate, quiet interview room, otherwise conversation will be inhibited and information lost. Posture and attitude convey sincerity, concern and how seriously problems are considered. The patient needs to form a trusting relationship, and a respectful, honest but never patronizing approach is normally accepted. A sympathetic hand can reassure and encourage an anxious or suspicious patient.

The importance of establishing a positive therapeutic relationship at this early stage cannot be overstated, as it can have far-reaching effects, not only for the openness of discussion but also for future compliance and prognosis¹².

History

The nature of psychotic symptoms, their form, content and course must be detailed. Late-onset schizophrenia may develop insidiously over months or a year or more^{2,3}, delusional depression over a few months, delirium over days and dementia over 1–2 years. The intensity of paranoid ideas and their effect on behaviour assist diagnosis and the evaluation of risk. Associated symptoms, particularly affective and cognitive, should then be elicited.

Current and past medical problems and their temporal relationship to the onset of paranoid symptoms must be clearly established, including visual or auditory failure. Aetiologically significant hearing loss in late-onset schizophrenia is typically of long duration, severe and due to bilateral middle ear disease, often originating in early life^{42–45}. Details of prescribed and non-prescribed drugs, dosages and recent alterations are essential.

Previous episodes of mental disorder should be confirmed from medical records, when past diagnoses and response to treatment may quickly clarify a diagnostic dilemma. Careful enquiry might uncover past episodes of untreated, self-limiting illness^{3,31} and changes in behaviour may date the onset of current problems.

Premorbid personality and behaviour are important because departures from these in old age usually signify the onset of a morbid process. Forty to fifty per cent of late-onset schizophrenics have schizoid or paranoid premorbid traits^{2,3} and the diagnosis of personality disorder depends on establishing a life-long attitude. Brenes Jette and Winnett⁴⁶ emphasized the interaction of narcissistic personality traits and the psychosocial consequences of ageing in their psychodynamic formulation of late-onset paranoid disorder.

The genetic loading of schizophrenia in old age is less marked than with younger patients but a positive family history is often found^{2,7,20,47}. Odd behaviour or suicide among family members may be discovered when formal psychiatric treatment is absent.

Current social circumstances and recent change are of relevance to aetiology and management. The schizophrenias of late life are particularly associated with social isolation, but rarely with precipitating life events^{2,3}. Paranoid patients frequently have poor socioeconomic status and multiple difficulties¹³ and social support has prognostic implications¹². Enquiring about alcohol and drug abuse must not be avoided for fear of offending a respectable elderly person. The elderly are not without vice and may be less inclined to confess it.

Mental State Examination

The detailed psychopathology of these conditions is described elsewhere and only points relevant to the process of mental state examination will be mentioned here.

It is important to ensure that the patient understands the terminology used to elicit abnormal experiences and that a common language is being used. Eliciting paranoid and psychotic symptoms can be difficult, but with tact and careful choice of words most patients will participate in an exchange of ideas about their experiences. This must be an unthreatening process for the patient and it is unwise to challenge or trivialize complaints at an early stage. A neutral position is advisable until a firm relationship is established, when complaints may be gradually reframed so that they can be viewed by the patient as problems that can be relieved, rather than immovable realities that are not amenable to therapeutic intervention. Suggesting that "it's all imagination" will be considered insulting and the patient's confidence will be lost.

Insight is rarely retained and patients may not volunteer experiences if they interpret questions as purely an enquiry into the state of their health. Patients have limited ability to accept the presence of illness or recognize psychiatric experiences as pathological⁴⁸. Needless to say, the mental state examination must be thorough.

Physical Examination

A complete physical examination should be performed routinely, with particular emphasis on neurological status and sensory function. The association between sensory impairment and late-onset schizophrenia^{44,49} makes attention to this area important and remediable conditions may be found. Visual impairment, particularly due to cataracts, is often found in association with delusions and deafness^{43,50} and visual hallucinations may be as much to do with ocular pathology as psychiatric diagnosis⁵¹. A particular form of acute, elaborate visual hallucinosis, the Charles-Bonnet syndrome, is usually related to eye disease or cerebral organic disorder⁵². Simple clinical interview and self-reporting seriously underestimate sensory impairment and more detailed ophthalmic and audiometric examination may be necessary^{53–55}. A simple battery of laboratory investigations is required for all patients, including haematology, biochemistry, thyroid function, urinalysis and chest radiography.

Advanced neuroradiological techniques promise much for the future but have limited clinical application at the present time. Some authors recommend the routine use of computed tomography (CT) and MRI. These procedures frequently reveal structural cerebral abnormalities, including increased ventricular size and periventricular and deep white matter hyperintensities^{56,57}, although their clinical significance is uncertain^{58,59} and they appear to bear little relationship to clinical state or outcome^{60,61}. These findings have no diagnostic value and the role of neuroimaging in clinical practice, at present, is to exclude specific intracranial pathology, particularly space-occupying lesions suggested by clinical examination. Similarly, non-specific electrophysiological abnormalities are common⁵ and the EEG will only be of value in a minority of cases.

Psychometric testing may provide a useful baseline measure that can be serially repeated when the possibility of dementia arises⁶². Psychometric testing certainly reveals cognitive deficits in late-onset schizophrenic patients, particularly affecting frontal lobe and memory function⁵⁶. These rarely signify dementia⁶⁰ and are more like the deficits found with early-onset schizophrenia than Alzheimer's disease⁵⁶. They do not correlate with severity of psychosis or other clinical parameters^{60,62}.

CONCLUSION

Most patients with late-onset schizophrenia or primary delusional disorders will be adequately and preferably managed from home^{12,63}. The need to admit to hospital seems to be declining⁶⁴ and may be determined as much by social and physical factors or treatment compliance as degree of psychopathology. For the patient requiring more than outpatient treatment, a day hospital can provide the necessary facilities for more intensive assessment of mental state, physical health and functional level.

The multifactorial contributions from ageing, physical disability, sensory impairment and social factors demand a multi-professional approach and all relevant disciplines must be available and involved⁸. The evaluation of these conditions requires clinical skill, rigorous attention to detail and an holistic approach. The accuracy of diagnosis and success of management will depend on the quality of initial assessment and if diagnostic doubts exist, treatment should be postponed until the situation becomes clear. Occasionally a diagnostic trial of treatment will be justified.

The paranoid disorders of old age are stimulating, complex, challenging clinical problems that encompass the breadth of psychiatry, medicine and social sciences and their assessment and management will continue to appeal to the enquiring clinical mind.

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Aetiology, Genetics and Risk Factors

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There is considerable controversy about the classification of the late-onset non-affective psychoses, and their relationship to “typical” early-onset schizophrenia. Kraepelin delineated “paraphrenia”, a late-onset delusional state with prominent hallucinations, as distinct from dementia praecox. Mayer performed a follow-up study of a number of patients diagnosed by Kraepelin as having paraphrenia, and found that a high proportion had a longitudinal course of illness similar to that of dementia praecox patients. This study challenged the distinction between paraphrenia and dementia praecox and, once the Bleulerian label “schizophrenia” had been widely adopted, “paraphrenia” became a neglected term.

The term was resurrected by Roth¹ as “late paraphrenia”, although he used it to describe a rather different group of patients, namely those manifesting a paranoid delusional state with prominent hallucinations for the first time in very late life. Confusingly, ICD-9 retained Roth’s category, relabelled it “paraphrenia”, and subsumed it under “paranoid states”; this has been dropped altogether from ICD-10. Furthermore, definitions of “late onset” range from over the age of 40 (after Feighner *et al.*²) to over 60 (after Roth¹). The 3rd edition of the American Psychiatric Association’s *Diagnostic and Statistical Manual* (DSM-III) set an age cut-off of 45 years for schizophrenia, so that later-onset paranoid states were labelled “atypical psychosis” or “paranoid (delusional) disorder”. The revised editions of the manual (DSM-III-R and DSM-IV) removed this age stipulation, and it is now widely believed that schizophrenia can manifest for the first time at any age.

These changing conceptions make comparison across studies difficult. In this chapter, we refer to studies where a 40 year or later cut-off is used. We employ the term “late paraphrenia” for those patients with an illness akin to Roth’s description, with first manifestation usually after the age of 60. Delusional (paranoid) disorder is not specifically addressed here.

GENETICS

It is generally accepted that schizophrenia tends to run in families. Gottesman and Shields³, pooling data from a number of family studies, calculated the average morbid risk for schizophrenia (broadly defined) in first-degree relatives of schizophrenia probands to be around 10%, compared with about 1% in the general population. Rates are generally lower when operational criteria such as those of DSM-III are applied, but the higher relative risk in first-degree relatives compared with the general population remains. Twin studies consistently report higher concordance rates for schizophrenia in monozygotic (MZ) compared with

dizygotic (DZ) twin pairs, suggesting that, in part at least, this familial aggregation is due to genetic factors. This conclusion is supported by adoption studies, which show that biological risk for schizophrenia is carried with the proband, irrespective of family environment (i.e. “adopted-away” offspring).

Few studies have specifically assessed familial aggregation in late-onset schizophrenia (see Table 91.1) and these all have methodological shortcomings. Authors have often failed to give a clear definition of illness in either probands or relatives; standardized diagnostic instruments have not been employed; and assessments have not been blind to proband diagnosis. Furthermore, proband numbers are mostly small, and it is often unclear exactly how many family members were assessed, and whether adjustments were made for those who were not. Also, because of the late onset of illness, many family members (siblings, parents) would have died and thus have been unavailable for interview; it is also difficult to know whether relatives had passed through the period of risk for illness-onset. Controls have rarely been employed, reliance being placed on results from independent studies of earlier-onset patients, which have again used different methodologies and diagnostic criteria.

Thus, the conclusions that can be drawn are limited. Overall, studies of “late paraphrenia” patients show rates of schizophrenia in siblings that are intermediate between those for siblings of early-onset probands and the general population. The discrepancy in rates quoted for parents is probably due to problems in assessing the parental generation. Those studies with an earlier age cut-off report higher risks for schizophrenia in relatives approaching those in the relatives of early-onset probands. However, these studies have used broad, non-specific definitions of schizophrenia, and have thus probably overestimated rates. The very high rate in the study of Huber and colleagues (see Table 91.1) is probably due to the misdiagnosis of patients with affective disorders, as they report no relatives with affective illness. The more recent study of Howard *et al.*⁴, who employed a controlled family interview methodology, is more sound. These researchers assessed lifetime risk of schizophrenia and other psychiatric disorders in 269 relatives of 47 late paraphrenia patients, and compared the rates with those in 272 relatives of 42 elderly general population controls. The results, shown in Table 91.1, reveal no difference in the risk between the two groups of relatives with respect to schizophrenia, but do show the relatives of the schizophrenia probands to have a significantly elevated risk of depression (16.3% vs. 4.4% in controls).

Kendler *et al.*⁵ in a review of family and twin studies, found no strong consistent relationship between age at onset of schizophrenia and risk for the illness in relatives. However, they did not specifically investigate late-onset schizophrenia patients, and the

Table 91.1 Family studies in late-onset schizophrenia

Reference No.	No.	Age	Definition of illness	Risk of schizophrenia	Risk of affective disorder
53 ^a	65	> 40	Schneiderian first rank	Siblings: 9.8% Parents: 4.4% (9 of 24 late-onset)	Not stated
25	57	> 60	"Late paraphrenia"	Siblings: 2.5% Parents: 0 Children: 7.3% Nephews/nieces: 3.1%	Not stated
14	93	> 60	"Late paraphrenia"	5 relatives (2 late-onset)	16 relatives
43	45	> 65	Systematized delusions with or without hallucinations	Siblings: 2.3% Fathers: 2.2% Mothers: 4.4%	Not stated
44 ^a	110	> 40	Schneiderian first rank	19.4%	0%
54 ^a	62 cases; 58 early-onset controls	> 50	Late- vs. early-onset schizophrenia (definition of illness not stated)	10.8% vs. 17.7% in early-onset patients	2.0% vs. 6.6% in early-onset patients
48	35 cases; 35 affective disorder controls	> 44	"Persistent delusions" with no affective features	5 relatives vs. 3 relatives of controls	0 relatives vs. 12 relatives of controls
4	47 cases; 42 controls	> 60	"Late paraphrenia" (cases) vs. non-psychiatric controls	Narrow age range (15–50 years): 1.3% in cases and controls; wide age range (15–90 years): 2.3% cases, 2.2% controls	21 relatives of cases vs. 6 relatives of controls ($p=0.003$)

^aQuoted by Volavka⁵⁵ (1985).
From ref. 56, with permission.

data reviewed here suggest that there may indeed be a gradient of familial risk, dependent upon age of onset of the proband. Such a notion is compatible with the idea that late-onset patients have a smaller genetic loading than those with early onset and thus require more "environmental events" to manifest the illness. This conception presumes a multifactorial "vulnerability/stress" or "continuum of liability" model of schizophrenia that is useful in as far as it produces testable hypotheses. For example, Holden⁶, in a study of 47 cases of paranoid psychosis with onset after the age of 60, found that deafness, which is considered a risk factor for late-onset schizophrenia (see below), was inversely related to family history of psychiatric illness.

An equally parsimonious explanation for the intermediate family risk in late-onset probands is that it is due to the greater proportion of such patients who have a less genetic subtype of illness. For example, paranoid ideation is common in late-onset schizophrenia patients, and Tsuang and Winokur⁷ and Tsuang *et al.*⁸ have reported that patients with the paranoid subtype of schizophrenia have later onset and fewer affected relatives than do the more disorganized "hebephrenia" patients. Furthermore, at least some patients labelled "late-onset schizophrenics" have an organic illness (see below), and such patients would be expected to have a low family risk for schizophrenia. Also, the family loading for affective disorder found in some studies (see above) suggests that at least some patients with late-onset schizophrenia have aetiological links with affective disorder.

FEMALE GENDER

All studies of late-onset schizophrenia that have included both sexes have attested to an excess of women. Table 91.2 shows data from a representative sample of such studies, which confirm that this female excess is robust to more- or less-restrictive definitions of illness. This excess cannot be explained on the basis of the relative longevity of women.

What is also evident from the data in Table 91.2, is that the older the sample, the greater the female excess. Few studies have assessed gender differences in non-affective psychoses in an epidemiologically-based sample of patients, ascertained across all ages at onset. The Camberwell Register First Episode Study⁹

afforded an opportunity to do this. The sample consisted of 91% of all patients ($n=477$) with a non-affective psychotic disorder, from a defined catchment area, who made their first contact with the psychiatric services over the period 1965–1984. Patients were rediagnosed according to a range of operational definitions for psychotic disorders, using the OPCRIT diagnostic system. Age-at-onset incidence curves were established for both sexes, using the base population figures as the denominator. Not only was the mean age at onset later for females, but the distributions of onset age for the two sexes were not isomorphic; males showed a dramatic early peak and a lesser mid-life peak, whilst females showed three peaks of onset, one in very late life. When subjected to an admixture analysis, the distributions for males showed two age distributions, with modal ages at onset of 21 and 39 years, whilst for females there were three distributions, with modal onsets at 22, 37 and 62 years¹⁰.

A possible explanation for the female excess in late-onset schizophrenia is that female schizophrenia patients are somehow "protected" from the manifestations of disease at earlier ages. For example, Seeman¹¹ has suggested that the antidopaminergic action of oestrogen has such a protective function, with the illness manifesting perimenopausally when oestrogen levels fall. This theory gains support from animal and clinical studies that show that oestrogen has antidopaminergic properties. However, as described above, the incidence curves for schizophrenia in women do not mirror the menopausal fall in oestrogen levels, and the disease can manifest for the first time at very advanced ages.

A further consideration in attempting to explain the female excess amongst late-onset schizophrenia patients is the fact that the brains of males and females age differently. Of particular interest is the differential rate of loss of dopamine D2 receptors, with loss being more precipitous in men than in women. Thus, young males have a relative excess of D2 receptors compared with females, but in older females this gender difference is reversed¹². This differential loss of D2 receptors between the sexes could conceivably play a part in the vulnerability of females to the manifestation of schizophrenia in late life and might also account, in part at least, for the particular female vulnerability to the development of tardive dyskinesia on exposure to neuroleptic medication in late life.

Table 91.2 Selected series of late-onset schizophrenia patients reporting gender ratio

Reference No.	No. of cases	Ascertainment method	Diagnosis	Age (years)	Ratio female:male
25	57	Hospital admissions	Late paraphrenia ^a	>60	5.3:1
43	47	Hospital admissions	Systematized delusion ± hallucinations; not demented	>65	22.5:1
44	644	Hospital admissions	Late-onset schizophrenia; not organic	>40	1.8:1
45	6064	First admissions	ICD-8 schizophrenia	>40	1.6:1
46	320	Hospital admissions	Late paraphrenia ^a	>65	6:1
47	25	Consecutive referrals	Late paraphrenia ^a	>60	3.2:1
48	35	Hospital admissions	Persistent delusional state; absence of mood or cognitive disorder	Onset >40	10.7:1
49	106	First admissions	ICD-8 schizophrenia, paranoid state, reactive psychosis, other psychoses	>60	2.2:1
6	37	Case register	Late paraphrenia ^a (13 cases considered “organic” at follow-up)	>60	7:1–3:1 ^b
50	477	Case register	ICD-9 schizophrenia and related disorders, paraphrenia, atypical psychoses	>60	4.4:1
51	47	Referrals from a number of psychiatric settings	Late paraphrenia ^a	>65	9:1

^aAkin to Roth's¹ criteria.

^bDependent on whether “organic” cases included.
From ref. 52, with permission.

PREMORBID CHARACTERISTICS

It has been consistently reported that a higher proportion of patients with late-onset paranoid psychoses have abnormal premorbid personality traits, most commonly described as “suspicious”, “hostile” and “reclusive”. Some workers have considered that the occurrence of paranoid psychosis in individuals with such personality traits is an “understandable transition”, while Retterstol¹³ has suggested that psychotic breakdown is a reaction to stress in a “hypersensitive” personality. Post¹⁴ proposed that the paranoid/schizoid personality traits reflect a long-standing latent schizophrenic disorder that manifests itself only when additional factors come into effect. Such factors might include social deprivation, sensory deprivation (deafness and possibly visual loss, as discussed below), frank cerebral pathology, or even ordinary ageing processes in the brain. Moreover, such personality traits could be expected to result in social isolation and reclusiveness, which in turn would exacerbate the paranoid imaginings characteristic of the illness.

In contrast to their early-onset counterparts, late-onset schizophrenia patients are generally neither educationally nor occupationally compromised. This again suggests that they may have a form of disease that is relatively distinct from the severe early-onset “dementia praecox” type.

SOCIAL ISOLATION

Ageing often results in increasing social isolation. It appears, however, that patients with late-onset schizophrenia have a greater likelihood of being socially isolated than age-matched normals or affective disorder patients. Low rates of marriage, few offspring and paranoid premorbid personality traits could all be expected to contribute to such isolation. Paranoid ideation in the disease itself is often directed at neighbours, resulting in further reclusiveness. Thus, social isolation in late-life psychosis might well be a consequence rather than a cause of the illness.

SENSORY DEFICITS

A number of general population studies have found an association between paranoid ideation and sensory impairment in the

elderly. This perhaps understandable association has also been investigated as potentially causal in late-onset paranoid illnesses. For example, Post¹⁴ reported that 25% of 72 elderly paranoid psychotic patients had hearing loss, compared with 11% of an affective disorder group; the mode of audiometric assessment was not stated. Cooper *et al.*¹⁵ found that 25 (46%) of a group of 65 elderly paranoid patients were “socially deaf”, compared with 12 (21%) of 67 patients with affective illness. Audiometry and otological examination of an enlarged sample (27 paranoid and 18 affective deaf subjects) revealed that the paranoid group were more likely to have long-standing conductive hearing loss, as opposed to later-onset sensorineural loss in the affective group. Visual impairment, mostly due to cataracts, was found in 30 (56%) of 54 of the paranoid group, and 21 (37%) of 57 of the affective group.

From such data, Kay *et al.*¹⁶ and Cooper¹⁷ concluded that in the elderly, long-standing conductive deafness is an independent risk factor for paranoid, as opposed to affective, psychosis. This probably does not apply in earlier-onset schizophrenia patients, neither are persons with profound (prelingual) deafness predisposed to schizophrenia. The association with visual impairment is less robust, possibly because determination of functional visual impairment is difficult.

More recently, however, Prager and Jeste¹⁸ failed to confirm an excess of constitutional (“uncorrected”) visual and hearing impairment in late-onset schizophrenia patients; they did, however, find that such patients were more likely than controls to have sensory deficits that were not adequately “corrected”, e.g. by spectacles or hearing aids. Thus, these data are complex and interpretation of causality problematic. Furthermore, there is no consistent association between particular modalities of sensory loss and any specific psychotic symptom. For example, it is not the case that hearing impairment is necessarily associated with auditory hallucinosis. Likewise, the association between visual impairment and visual hallucinations, so dramatically represented in the Charles–Bonnet syndrome¹⁹, does not appear to be found in late-life schizophrenia.

The social and psychological consequences of sensory impairments (social withdrawal and ostracization; misinterpretation of social cues) may result in suspiciousness and hostility, leading to paranoid ideation. Abnormal percepts associated with reduced sensory input might result in hallucinations, as in the Charles–Bonnet syndrome, although secondary delusional elaboration is

considered unusual in such settings¹⁹. Moreover, Watt²⁰, in a study of 35 patients, found no association between hearing loss and paranoid psychosis manifesting in middle life. It might be that psychosis occurs only in individuals who are already prone to paranoid ideation, e.g. by virtue of mild cerebral damage. This is consonant with findings in an elderly general population sample²¹, of an association between persecutory ideation, sensory impairment and cognitive dysfunction.

“ORGANIC” FACTORS

Acute paranoid ideation may result from cerebral or extracerebral organic factors, and the elderly are especially susceptible to such effects. More persistent delusional persecutory states are seen in association with a wide variety of structural brain changes and systemic toxic and metabolic disturbances. In particular, such states can occur in dementia. For example, Wragg and Jest²², in a review of studies reporting the prevalence of psychotic phenomena in patients with Alzheimer’s disease, found a rate of delusions of anything from 10% to 73% (aggregating to 30–38%, with a median of 33.5%). For hallucinations, the range was 21–49% (mean 28%). The importance of organic factors in the aetiology of psychotic phenomena in the elderly is underlined by general population studies, which have consistently found the strongest predictor of paranoid ideation in the elderly to be cognitive impairment^{23,24}.

Kay²⁵ stated that a diagnosis of late-onset schizophrenia can be made only in the absence of “gross and persistent disorientation in time and place or severe failure of memory”. Roth¹ claimed that his follow-up data validated the separation of such disorders from the dementias, while Kay and Roth²⁶ found only minimal evidence of organic cerebral damage in their late paraphrenic patients. Subsequently, Kay²⁵ reported survival rates for late paraphrenia patients of 0.97:1 (observed expected), compared with 0.3:1 for patients with dementia.

However, it is becoming increasingly clear that a significant number of elderly patients with “functional” paranoid states *do* have evidence of organic cerebral deterioration, not part of ordinary ageing. For example, Post¹⁴ reported that some 15% of his patients were demented at follow-up, while Holden⁶ found that 13 (35%) of 37 late paraphrenia patients had demented within 3 years of diagnosis; the only distinguishing feature on admission was a slight impairment on psychometric assessment in the “organic” group.

Neuroimaging investigations have revealed structural brain changes in a proportion of elderly patients with persistent psychosis and no obvious neurological or neuropsychological deficit. Miller and colleagues²⁷ reported three such patients who had CT scan evidence of cerebral infarction and one with normal pressure hydrocephalus. In a prospective CT and MRI study of 27 patients with late-life psychosis, Miller *et al.*²⁸ found silent vascular lesions in five (19%); subcortical frontal connections were most commonly involved. Similarly, Jernigan *et al.*²⁹ reported 13 patients with late-onset psychosis (10 schizophrenia, three delusional disorder) to have significantly more white-matter pathology on MRI than age-matched normal controls; and Breitner *et al.*³⁰ found significant leucoencephalopathy, especially affecting the temporoparietal and occipital regions, in eight late-onset schizophrenia patients; such lesions were minimal or absent in controls.

These findings may provide useful information about the pathogenesis of late-onset schizophrenia. However, the exact relationship of the reported lesions to the manifestation of the disease is unclear. There is little consistency in the site of the lesions, and white-matter changes have also been reported in elderly patients with severe depression³¹. Moreover, it appears

that when late-onset schizophrenia patients are clinically screened fastidiously, so as to exclude any patients with potential “organic” aetiologies, they show no excess of white matter abnormalities compared to controls³².

NON-SPECIFIC STRUCTURAL BRAIN ABNORMALITIES

Numerous neuroimaging studies have revealed an increase in ventricular:brain ratio (VBR) in younger schizophrenia patients compared with normal controls. These findings, and those suggesting temporal lobe dysplasia, appear to have a developmental origin. There are few such studies in late-onset patients. Rabins *et al.*³³ studied 29 schizophrenia patients with disease onset after the age of 44, and found mean VBR to be greater than for matched normal controls. Naguib and Levy³⁴ reported similar findings in 43 late paraphrenia patients; there was no correlation between illness duration and ventricular size, and ventricular size did not predict disease outcome at a mean of 3.7 years³⁵. In an MRI study, Jeste and Harris³⁶ confirmed an increase in ventricular size in 20 late-onset schizophrenia patients compared with normals. Again, there was no relationship between illness duration and ventricular size, suggesting that the abnormality preceded disease onset and was not progressive. In an attempt to correlate brain imaging findings with clinical symptomatology, Howard and colleagues³⁷ reported that ventricular/sulcal enlargement was most dramatically evident in those late paraphrenia patients who did not manifest Schneiderian first-rank symptoms.

The relationship between neuroimaging findings in early- vs. late-onset schizophrenia patients remains a moot point. Pearlson *et al.*³⁸ compared 11 late-onset with 11 early-onset schizophrenia patients, and found no significant differences between the groups in terms of volume of a number of brain structures. Summarizing these and other such data, Pearlson³⁹ concluded that “early- and late-onset cases of schizophrenia share common structural ... brain abnormalities”. But the significance of these findings is unclear. It seems very unlikely that they have a similar developmental origin to the equivalent abnormalities in early-onset schizophrenia. Indeed, a comparative study of very early (<25 years) and very late-onset (>60 years) schizophrenia patients found that the latter group did not seem prone to those risk factors (e.g. obstetric complications) that have been implicated in the aetiology of the neurodevelopmental form of schizophrenia⁴⁰. Presumably, the ventricular enlargement is of little consequence in its own right, being merely an echo of some undetected cerebral lesion. Burns *et al.*⁴¹ have suggested that there occurs, in late paraphrenia, an “uncoupling” of the normal association between ventricular and cortical size, but the exact mechanism whereby this results in psychotic symptoms is unclear. There is a place for further studies in this area. Specifically, further attempts should be made to find clinical correlates of enlarged ventricles, and to assess predictive value in longer-term follow-up studies. The use of more advanced neuroimaging techniques to search for more specific abnormalities will also be important.

CONCLUSIONS

Many authors consider late-onset schizophrenia to be “a form of schizophrenia, albeit attenuated and modified”. One suggestion is that inherent genetic vulnerability, itself insufficient to cause psychosis, acts in concert with environmental factors (social isolation, sensory impairments, non-specific brain atrophy) to precipitate delusional breakdown. Paranoid/schizoid premorbid personality traits might be an expression of such intermediate genetic loading.

Alternatively, the late-onset schizophrenia-like states could be considered as comprising one or more subtypes, relatively distinct from early-onset schizophrenia. For example, Post¹⁴ delineated “paranoid psychosis”, “schizophreniform psychosis” and “late schizophrenia”; however, this schema has neither prognostic nor aetiological utility. A more useful approach might be subdivision based on biological parameters that have a bearing on pathogenesis, e.g. the finding of a subgroup of patients with subcortical white-matter changes has important heuristic implications.

Our own view is conditioned by our belief that early-onset schizophrenia is a neurodevelopmental disorder whose origins lie in faulty brain development in foetal or neonatal life⁴². It seems a long bow to draw to suggest that patients with the first manifestation of psychosis in late or very late life have a neurodevelopmental illness. In this context, we consider that those patients currently labelled “late-onset schizophrenia” form a heterogeneous group, some of whom have an illness relating to paranoid personality, some to sensory deprivation, some to late-onset organic change and some to affective illness. A degree of interaction will be expected between these factors. Careful clinical, neuroimaging and, ultimately, pathological examination of such patients will be required to further elucidate these issues.

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Brain Imaging in Schizophrenia-like and Paranoid Disorders in Late Life

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Two general conclusions can be drawn from the results of brain-imaging investigations of patients with an onset in late life of a schizophrenia-like psychosis. First, the established neuroimaging abnormalities reported in early adult life-onset schizophrenia are also seen in later life-onset patients. Second, imaging has not implicated focal or generalized neurodegenerative abnormalities that can be implicated in psychosis aetiology, as has been the case for late-onset affective disorders.

STRUCTURAL IMAGING

Computed tomography (CT) studies of onset after 45 years¹ and 60 years² patients have shown lateral and third ventricle volume enlargement that does not approach the ventriculomegaly seen in Alzheimer's disease. Structural magnetic resonance imaging (MRI) has confirmed these findings, as well as indicating possible reductions in left temporal lobe and superior temporal gyrus volumes^{3–5}. MRI studies of patients with so-called "late-life psychosis" have included patients with organic as well as schizophrenia-like psychoses and, not surprisingly, have reported an excess of focal structural abnormalities within deep grey and white matter structures⁶. Similar studies of patients with more rigorously defined late-onset schizophrenia, with exclusion of individuals who are cognitively impaired or have clinical evidence of focal cerebrovascular disease, have found no significant increase in such focal brain lesions compared to age-matched comparison subjects^{7,8}.

FUNCTIONAL IMAGING

Resting bloodflow abnormalities have been reported with single-photon emission computed tomography (SPECT) in "late-life psychosis", which may include patients with organic psychoses⁹. To date no bloodflow studies using cognitive activation paradigms have been reported in late-onset patients. A single neuroreceptor positron emission tomography study has reported increased binding values for dopamine D2 receptors³, although

age-matched control subjects were not examined for comparison. In a small group of drug-naive onset 60+ patients, we were unable to demonstrate any absolute increase in striatal D2 receptor number compared with elderly controls using SPECT¹⁰. Hence, the issue as to whether or not neuroreceptor levels are abnormal in late-onset cases mirrors the dispute in the early-onset schizophrenia literature.

Although brain imaging studies to date have supported the concept that early- and late-onset cases of schizophrenia share common structural and neuroreceptor brain abnormalities, more sophisticated future studies may show differences. These may help to settle which of two current hypotheses concerning the aetiopathology of late-onset schizophrenia and very late-onset schizophrenia-like psychosis are most likely to be correct. If schizophrenia arising at any point in life is essentially a unitary condition, then we cannot expect neuroimaging studies of even remarkable sophistication to reveal differences between early- and late-onset cases¹¹. If (and the author favours this second hypothesis) the later-onset cases represent a subtle organic phenocopy of schizophrenia, then application of novel imaging methodologies, such as diffusion tensor imaging, which allow high resolution definition of white matter structures *in vivo* should be informative in the next few years.

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Although brain imaging studies to date have supported the concept that early- and late-onset cases of schizophrenia share common structural and neuroreceptor brain abnormalities, more sophisticated future studies may show differences. These may help to settle which of two current hypotheses concerning the aetiopathology of late-onset schizophrenia and very late-onset schizophrenia-like psychosis are most likely to be correct. If schizophrenia arising at any point in life is essentially a unitary condition, then we cannot expect neuroimaging studies of even remarkable sophistication to reveal differences between early- and late-onset cases¹¹. If (and the author favours this second hypothesis) the later-onset cases represent a subtle organic phenocopy of schizophrenia, then application of novel imaging methodologies, such as diffusion tensor imaging, which allow high resolution definition of white matter structures *in vivo* should be informative in the next few years.

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Schizophrenic Disorder and Mood-incongruent Paranoid States: Epidemiology, Prevalence, Incidence and Course

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Although paranoid symptoms are commonly encountered in a range of psychiatric disorders that present to old age psychiatrists, remarkably few epidemiological studies of mood-incongruent paranoid states have been reported compared with cognitive and affective disorders¹. Because these conditions are comparatively rare, it is difficult to determine their prevalence and incidence accurately. The epidemiological information that we do have on such states that have their onset in later life has come from two sources: studies based upon patients in contact with psychiatric services, and surveys of the elderly general population. Because of the nature of paranoid states, sufferers are unlikely to cooperate with community surveys and are often hidden from contact with psychiatric services. Thus, all the studies reviewed below are likely to represent underestimates of the true prevalence and incidence of these disorders.

PARANOID IDEATION AMONG COMMUNITY-LIVING ELDERLY

In 1173 subjects aged over 64 years in the Epidemiological Catchment Area survey, with a response rate of 85%, generalized persecutory ideation, as assessed on the paranoid scale of the Mini-Mult, was present in 4%². There was a significant excess of unmarried individuals among those with persecutory ideation, but no association with gender or living alone. Other associated features were visual and hearing deficits, cognitive impairment, impaired physical health and disabilities in daily living, together with reduced social and economic resources. Among a community sample of 1420 individuals aged over 75 years, paranoid ideation (defined as recording of paranoid symptoms by both a physician and informant's interview) was found in 6.3%³. The prevalence of paranoid ideation in people with cognitive impairment was 12.1%, while it was only 2.6% in those who were cognitively intact. Once the effect of cognitive impairment has been controlled for, the variables significantly associated with paranoid ideation were being divorced, being female, having depressive symptoms, receiving psychotropic drugs, having no friends or visitors, using community care and being an immigrant. In a survey of 935 interviews with individuals aged 70 or over in Canberra and a neighbouring town⁴, 65 had at least one psychotic symptom; 22 reported current auditory hallucinations only, 23 delusions only and three hallucinations and delusions: 25 of these individuals had cognitive impairment or dementia. The point prevalence of

psychotic symptoms was 5.7% and the significantly associated risk factors, apart from cognitive impairment, were living alone, male gender, limited education, social isolation, poor health and depressive symptoms.

HALLUCINATIONS IN COMMUNITY-LIVING ELDERLY

Data from the Epidemiological Catchment Area study were used to estimate the self-reported age-specific prevalence of hallucinations in a sample of 15 258 individuals of all ages. Patients with dementia were not excluded, and although the prevalence of both auditory and visual hallucinations was highest in young subjects, an increase in auditory and visual hallucinations was found in the elderly, with a rate of visual hallucinations of 40/1000/year in males aged 80+⁵.

PSYCHOSIS DIAGNOSED IN COMMUNITY-LIVING ELDERLY

The prevalence of schizophrenia and delusional disorder in those aged 65+ in the community has varied widely from study to study, but on the whole low rates have been found. The Epidemiological Catchment Area survey found a prevalence of schizophrenia of 0.2%⁶ and the 6 month prevalence rate of schizophrenia was 0.4–0.6% in a Danish survey⁷. In a sample of 612 elderly Chinese Singaporeans examined using GMS–AGECAT criteria, 0.5% had schizophrenia or paraphrenia diagnoses⁸. From a random sample of 5222 individuals aged 65+, Copeland and colleagues⁹ made estimates of the prevalence and incidence of DSM-III-R-defined delusional disorder and schizophrenia. The sample were chosen from the lists of general practitioners' patients and were interviewed by nurses trained in the use of the GMS–AGECAT computerized diagnostic system. The prevalence of DSM-III-R schizophrenia was estimated at 0.12% (95% CI, 0.04–0.25%) and delusional disorder at 0.04% (95% CI, 0.00–0.14%). The minimum incidence of schizophrenia for new cases was 3.0, for new and relapsed cases 45.0, and for delusional disorder 15.6/100 000/year. Two of the five cases of schizophrenia identified in the sample were found to have been first diagnosed before the age of 65.

STUDIES BASED ON PSYCHIATRIC CONTACTS

From the 1966 official figures from England and Wales for individuals aged over 65 years, Kay¹⁰ calculated annual incidence rates of schizophrenia of 10–15/100 000 for males and 20–25/100 000 for females. The annual incidence of DSM-III-R-defined schizophrenia in the over-65s on the Camberwell Case Register was estimated at 12.6 per 100 000¹¹. van Os and colleagues¹² examined the annual incidence rate of late-onset (age 59+) non-affective, non-organic psychosis in 8010 elderly admissions to psychiatric hospitals in The Netherlands and 1777 elderly admissions in the UK. The incidence of psychosis showed a significant increase with age in both countries, the cases rising from around 10/100 000 person-years in the age group 60–65 years to just over 25/100 000 person-years in the 90+ age group. After adjustments for the possible confounding effects of time trend and gender, the linear trend in the association between increasing age and first admission rates corresponded to an 11% increase in incidence with each 5 year increase in age.

COURSE AND COGNITIVE PROGNOSIS

Long-term follow-up of those patients we used to call “late paraphrenics”, but should now describe as suffering from “very late-onset schizophrenia-like psychosis”¹³, has shown that in the absence of cognitive impairment at the outset, the mortality rate does not differ from expectation and the causes of death in these patients are similar to the general population¹⁴. When those patients with accompanying organic brain syndromes and associated cognitive impairment are included in follow-up studies, rates of progression to dementia and mortality are, not surprisingly, high¹⁵.

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The Fate of Schizophrenia with Advancing Age: Research Findings and Implications for Clinical Care

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Although approximately 20% of the long-stay population of psychiatric hospitals in the UK are aged over 65 years¹, and the large-scale closure of mental hospitals that began in the 1950s is nearing completion, old schizophrenic patients and the factors that might influence the success of their move into the community have only recently become the subject of serious study². Ready availability of nursing-home places and a lack of public concern about the potential dangerousness of elderly people with schizophrenia in the community have fostered such disinterest, although the numbers of people involved are enormous: some 200 000 elderly people with schizophrenia have been discharged to nursing homes in the USA³.

LONG-TERM OUTCOME IN SCHIZOPHRENIA

Enduring positive symptoms of psychosis, cognitive deficits, negative symptoms and adaptive problems may all contribute to long-term disability in schizophrenia. Although some long-term follow-up studies suggest that cognitive and social decline plateau after the age of 40 years and that some symptoms actually improve with increasing age⁴⁻⁶, many patients experience multiple exacerbations of illness or spend many years in hospital⁷. Although we are traditionally taught that males have a generally worse prognosis in schizophrenia, the situation is complicated. At 2–13 years after initial diagnosis, females do seem to do better^{8,9}, but their advantage may be lost after 20 years^{4,6}. From follow-up studies spanning three decades, Seeman¹⁰ has suggested that males are typically more severely ill in the first 10 years of illness but then improve, while females, after a relatively need-free first decade, show increasing disability over time.

COGNITIVE IMPAIRMENT

The mean Mini-Mental State Examination (MMSE) score in a group of 38 chronic elderly schizophrenic inpatients in one study was 9.6¹¹—well within the moderately demented range. Such cognitive deficit does not appear to be specifically linked to a history of brain-damaging treatments, such as leucotomy, insulin coma therapy or lifetime total neuroleptic dosing⁷ and cannot be attributed to the appearance of Alzheimer-type neuropathological change, which is rare in the brains of such patients coming to neuropathology¹². Indeed, in a neuropathological study of 66 patients with schizophrenia, 68% of whom had a history of marked cognitive impairment in life, only 8% had brain changes

that satisfied diagnostic criteria for AD¹³. Although cognitive impairment in such patients cannot therefore be attributable to AD, those patients with most severe impairment had significantly more plaques and tangles in their brains than did the unimpaired subjects examined. Severe cognitive impairment is the most important single predictor of poor outcome in chronic schizophrenia¹⁴. There is debate in the literature regarding whether or not schizophrenic cognitive impairment progresses within chronic illness. Although cross-sectional studies have tended to suggest that cognitive function does not decline over time in schizophrenia^{15,16}, there is no doubt that cognitive impairment does appear at some point in the illness. Either it is present as a static feature at or soon after illness onset, or it progresses insidiously or in limited subgroups of patients, so that the studies have missed it. This latter point is supported by the results of a large 30-month cognitive follow-up study of 326 chronic schizophrenic patients aged over 65 years¹⁷. In this study, 30% of patients who had baseline scores in the less impaired range of the Clinical Dementia Rating scale showed a worsening of this score at follow-up to moderate or more severely impaired levels. Only 7% of the sample with lower scores at baseline showed any improvement in functioning. Factors predicting cognitive decline included lower levels of education, older age and more severe positive symptoms. Although the cognitive and functional decline over 30 months demonstrated by this study was convincing and dramatic, it is important to remember that the patients involved were chronically institutionalized, with very adverse illness courses. Middle-aged poor-prognosis schizophrenic patients may also have neuroimaging evidence of progressive ventricular enlargement¹⁸—further support for the thesis that a subgroup of middle-aged or elderly schizophrenic patients with poorly controlled symptoms are at particular risk of declining cognitive function. Although cognitive deficits have received the most research attention, because they are relatively straightforward to measure reliably and appear to be so predictive of outcome, they represent only a single dimension of disability in chronic schizophrenia. In a study of 102 middle-aged or elderly outpatients with schizophrenia, assessed with the Direct Assessment of Functional Status (a standardized measure of behaviours during performance of simulated daily tasks), patients were significantly more limited than controls on all subscales, except for grooming and eating¹⁹. Schizophrenic patients were more disabled than outpatients with major depression, but less impaired than those with Alzheimer's disease. Lower levels of formal education, greater severity of extrapyramidal symptoms and cognitive deficits were all associated with lower functional assessment scores.

THE EFFECTS OF INSTITUTIONALIZATION

Since poor long-term outcome in schizophrenia is associated with long institutionalization and cognitive impairments, could the deficits be a consequence of a non-stimulating psychiatric hospital environment or overtreatment with medication? One simple way to examine the possible causal relationship between these factors has involved comparing the cognitive and adaptive functioning deficits in groups of elderly schizophrenic patients cared for in different types of institutions in widely differing parts of the world. In a cross-national study of cognitive impairment in poor-outcome geriatric patients with schizophrenia²⁰ in London and New York, remarkable similarities in cognitive dysfunction between the US and UK patients were found, despite differences in the structure of institutional care provided. Mean MMSE scores in New York were 10.5 and in London 10.6. When the patients in the two centres were investigated with an adaptive functioning scale, however, differences emerged which were probably related to institutional differences. American patients were more impaired in social initiation, but less impaired in social competence and personal hygiene, than their English counterparts.

WHERE ARE PATIENTS BEST LOOKED AFTER?

It is probably unsafe to make generalized assertions about the ways in which the needs of elderly schizophrenic patients are best met because they are a heterogeneous group in terms of enduring psychotic symptoms, cognitive and adaptive deficits and family or other social support. In a cross-sectional study of 97 chronically hospitalized schizophrenic patients, 37 chronic schizophrenic residents in nursing homes and 31 acutely admitted geriatric patients with schizophrenia, patients in each of these groups had very different patterns of symptoms and impairments²¹. Whilst differences in positive and negative schizophrenic symptoms were small, nursing-home residents had the most severe adaptive deficits. Prospective studies of chronic schizophrenic patients successfully discharged to nursing homes, compared with those who are retained in long-term psychiatric care, have shown that it is not cognitive or adaptive deficits that prevent discharge but continuing belligerence and hostility²². Just as it appears unwise to generalize about the care needs of patients, it is rash to assume that all nursing homes or long-stay psychiatric facilities are the same. When 159 long-term schizophrenic inpatients within Veterans Administration hospitals were allocated to either a community nursing home, a Veterans Administration nursing home or another long-stay psychiatric ward, or allowed to remain on their original ward, at 12 months the patients with the best outcomes were those who had been transferred to another long-stay ward. The worst outcomes were seen in those who had been discharged to community nursing homes^{23,24}. Rather than the location of care, particular features of the quality of care were the factors most strongly associated with good outcome. Staffing characteristics, e.g. the staff:patient ratio and the rate of staff turnover, together with the mean functional ability of fellow residents, were significantly linked to outcome. A similar study from the UK came to superficially conflicting conclusions. Elderly long-stay schizophrenic patients transferred to nursing homes showed slower functional decline over the next 2–3 years than those who remained on the wards²⁵. The important difference from the situation in the Veterans Administration study was that in the UK study staff–patient contact was greater in community facilities than on the long-stay wards and this contributed to better outcome. Studies such as those reviewed above examine only the grosser disabilities and deficits of elderly schizophrenic patients and the few published investigations of quality of life give

a rather depressing insight into exactly what the lives of these people are like. In a 10 year follow-up of 40 older patients with schizophrenia, overall subjective quality-of-life ratings did not improve from the low levels seen at the beginning of the study²⁶. Ratings on a small number of items (contacts, inner experiences and knowledge/education) had improved slightly, but the reasons for these improvements were just as likely to be that patients had downgraded their expectations as that they were interacting more successfully with their environment or that housekeeping services received had improved.

MORTALITY IN CHRONIC SCHIZOPHRENIA

Excess mortality among patients with schizophrenia is a consistent and accepted research finding that cannot be fully explained by the observation that 10% of patients will kill themselves²⁷. In a prospective study of 88 schizophrenic patients with a mean age of 62.6 years at study commencement, 39 had died after 10 years' follow-up—none through suicide²⁸. The relative risk of death among the patients was 1.33 (95% CI, 1.01–1.65). Six variables, some of which have important clinical implications, affected independent prediction of reduced survival and these were: increasing age; male gender; the edentulous state; time since most recent neuroleptic withdrawal; maximum number of antipsychotics given concurrently; and the absence of anticholinergic treatment.

IMPLICATIONS FOR CLINICAL CARE

Since elderly patients with chronic schizophrenia have high levels of disability and dependence upon caring services, their apparent invisibility to mental health policy makers must largely be attributable to ageist assumptions that they should not expect much more than basic nursing home provision, and to a lack of general public concern about the safety of placing elderly (and therefore “low-risk”) psychotic patients in the community. They also represent a group of patients who are poorly served by specialist psychiatric services. Sometimes rather grandly termed “graduates” because they are alumni of mental health services set up for younger patients, it is often not clear whether they have become the responsibility of local old age psychiatrists or whether they should continue to be looked after by the general adult psychiatry teams with whom they have been in contact. In a call to arms to all mental health professionals who may be involved in the care of these patients, Rodríguez-Ferrer and Vassilas have set out four objectives of importance in the establishment of a seamless and ideal service. First, general practitioners should be central in the coordination of service provision and should be involved in the assessment of physical needs, as well as psychiatric ones. Second, the organization and delivery of specialist mental health services should take into account the fact that, in the future, the majority of these patients will live in residential and nursing homes. Third, purchasers of mental health services need to be aware of the effects of the quality of the physical and staffing environment of residential and nursing homes on patient functioning. Finally, services should maintain clarity at all times as to exactly which agencies (psychiatric, social, voluntary) have responsibility for each individual's care. Now that we have an agenda for the management of this hitherto-neglected group, together with novel antipsychotic agents that are less likely to induce movement disorders and may even improve cognitive function²⁹, this really does represent a clinical population for whom recent research has positive implications. The most recent indications are that deinstitutionalization has been, on the whole, a modest success. At the end of a 5 year follow-up of 670 elderly

chronic patients (80% of whom had schizophrenia), 89.6% were still living in the community and only one-third had required any kind of hospital readmission³⁰.

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Rehabilitation and Long-term Management

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THE LONG-TERM OUTCOME OF SCHIZOPHRENIA

Schizophrenia has been conventionally considered to be a disorder with a poor long-term prognosis. The concept of schizophrenia as a disorder with a poor prognosis was reflected by DSM-III¹, where features included a failure to return to pre-morbid functioning, recurrent acute exacerbation with increasing residual impairment between episodes, continued symptoms, unemployment, social isolation and an inability for self-care. DSM-IV² included a revised view of schizophrenia, stating that most studies of the outcome of schizophrenia suggest that the course may be variable, with some individuals displaying exacerbation and remission, whereas others remain chronically ill.

The pessimistic views on outcome had mainly been formed from the early longitudinal studies reported by Kraepelin³ and Eugen Bleuler⁴. In the course of his work, Kraepelin became pessimistic about the outcome of dementia praecox, reporting that fewer than 5% of patients sustained a lasting recovery. When Eugen Bleuler originally described the concept of schizophrenia, an essential component of the diagnosis was that patients were not able to regain their premorbid abilities. These early studies have had a major effect on our concept of schizophrenia. Indeed, many psychiatrists^{5,6} considered that if a patient makes a good recovery, then the validity of the diagnosis of schizophrenia may be in question. These views still persist and have had a major effect on service planning for people suffering from schizophrenia and their medical management. Five, large, long-term studies of schizophrenia patients have produced very different results from those of Kraepelin and Bleuler and have effectively challenged this view.

Harding *et al.*^{7,8} carried out a 32 year follow-up study of a rehabilitation programme of comprehensive rehabilitation and community placement for those "backward" patients who had not improved sufficiently following the introduction of chlorpromazine. A total of 269 patients who were considered to be amongst the most severely disabled and chronically mentally ill in the hospital were referred to the programme. After an average of 32 years, 178 were still alive and were interviewed again; 71 patients had died and in these cases the family were interviewed; 13 other patients were still alive but refused participation; and the remaining seven could not be located. A battery of interview instruments described as the Vermont Community Questionnaire was used to evaluate outcome. Evidence of good inter-rater reliability and face and construct validity for the questionnaires are presented in their paper. The patients were rediagnosed using DSM-III classification retrospectively: 188 of the original subjects were considered to have met the DSM-III criteria for schizophrenia. The results showed that the long-term outcome was mixed, with many patients demonstrating various degrees of

productivity, social involvements and competent functioning; 68% of the patients displayed neither positive nor negative symptoms of schizophrenia at follow-up and 45% of the sample displayed no psychiatric symptoms at all; 82% of patients had not been in hospital in the last year; 61% met with friends every week or two; 68% had one or more moderately/very close friends; 40% had been employed in the past year; 81% were rated as being able to meet basic needs; and 73% were felt to lead a moderate to very full life. The study also includes an assessment of income; 77% of the sample were assessed on the Community Care Schedule as having an adequate income. The schedule's definition of "adequate" was that "the amount of money received will cover the subject's basic needs comfortably". Four other studies report similar findings.

Manfred Bleuler⁹ described a follow-up study of 208 patients who met his own and Eugen Bleuler's diagnostic criteria for schizophrenia, and reported a significant improvement in 53% at a 23 year follow-up.

Huber *et al.*¹⁰ described the largest study, following up 502 patients over an average of 22.4 years, using the diagnostic criteria of E. and M. Bleuler^{4,9,11} and Schneider¹²; 53% showed significant or total improvement.

Ciompi and Muller¹³ followed up 289 patients admitted before the age of 65 who were over 65 by 1963. The mean length of follow-up was 36.9 years. The diagnostic criteria of E. and M. Bleuler were used; again, over 50% (57%) of patients showed significant improvement.

Tsuang *et al.*¹⁴ followed 186 patients admitted to Iowa State Hospital between 1934 and 1944 for a period of 35 years. In this study, 46% showed significant improvement.

The results of these studies challenge the prevailing concept of outcome in schizophrenia and support the concept of heterogeneity of outcome. Together, these studies found that between one-half and two-thirds of more than 1300 patients studied for a period in excess of 20 years achieved recovery or significant improvement. This is not to deny the fact that many patients do not show an improvement over the years and require continuing medical and social support, but it clearly suggests that it is erroneous to view deterioration as an inevitable outcome.

The differences between these studies and those of Kraepelin and Bleuler have been attributed to sampling biases. The criticism has been that both Kraepelin and Bleuler chose samples of patients who were admitted to hospital for long-term care. In addition to this, Kraepelin's studies probably included patients with tertiary syphilis and other organic disorders for which tests were unavailable at the time. Certainly, Eugen Bleuler's son, Manfred Bleuler, attributed his father's pessimism about the outcome of schizophrenia to this bias.

ENVIRONMENTAL, SOCIAL AND CULTURAL FACTORS AND OUTCOME OF SCHIZOPHRENIA

There is evidence that environmental factors play a major part in the outcome of schizophrenia, although this is clearly not as marked as was first claimed in the early descriptions of institutionalization by Barton¹⁵ and Goffman¹⁶. They considered that the disabilities of long-stay patients were primarily due to the psychiatric institutions in which they lived. In a major study of these institutions, Wing¹⁷ studied the outcome of patients who were managed in three hospitals with very different policies of care. He concluded that institutionalization had contributed significantly to the patients' disabilities.

The influence of environment and culture was also illustrated by the WHO international pilot study of schizophrenia¹⁸, in which 1202 patients diagnosed as having schizophrenia in 10 different countries were followed up over a 2 year period. The heterogeneity of outcome was again underlined, but patients suffering from schizophrenia in the developing countries had a better outcome than those in the developed countries. The report concluded that the diagnosis of schizophrenia alone did not provide sufficient grounds for a firm statement about the patient's likely pattern or course, probability of relapses and admissions, or the degree of social impairment in the future.

Similar findings emerge from a 5 year follow-up study of patients living in the peasant society of Sri Lanka, described by Waxler¹⁹. She concluded that in Western society, expectation and a belief about mental illness and the process by which treatment is provided alienate patients suffering from schizophrenia from their normal roles, and thus prolong illness. In contrast, the beliefs and practices in non-industrial societies encourage a quick return to normality.

THE ROLE OF REHABILITATION

Although the recent outcome studies have shown that the prognosis of schizophrenia is not as poor as had previously been considered, it is also clear that outcome is mixed, with some patients showing no improvement and remaining severely disabled. Continuing care services for people suffering from schizophrenia should aim to reduce these disabilities wherever possible and to meet the needs of those who remain severely disabled into old age.

The care of patients with schizophrenia has moved from long-term hospital to care in the community. This international change occurred with very little contemporary evidence to support it. However, the controlled studies that exist^{20,21} demonstrate that patients treated in the community did not experience increases in homelessness, mortality or suicide and did not need excessive readmissions to hospital. Importantly, patients taking part in these studies preferred treatment outside hospital.

Intercultural studies would support the concept that environmental factors can have an effect on outcome. However, although one would expect that the socioenvironmental factors involved in community care would improve the prognosis, this has been very difficult to demonstrate in practice. Wing's Three Hospital Study is one of the few papers that supports this view. There is little evidence that psychiatric symptomatology or psychosocial function improve when patients are treated in the community.

In the UK the view that all patients with schizophrenia can have long-term care in the community has been revised, and there is provision for 24 hour nursing care²². Treiman²³ demonstrated that this care can be successful in reducing severely disrupted behaviours. However, this group of patients are different from the long-stay hospital population. They are younger, suffer from more psychotic symptoms and often have multiple problems, such as

cognitive deficits, substance misuse and violence. Their self-care skills are often intact.

CHANGES IN REHABILITATION

One problem in interpreting these studies is that the term "rehabilitation" may mean very different things to different people, and indeed, in recent years the term "rehabilitation" has sometimes even become synonymous with discharge from hospital. This use of the word is misleading. Bennett²⁴ considered that the goal of rehabilitation was to enable the individual "to make the best use of his residual abilities in order to function at an optimum level in as normal a social context as possible". He introduced the concept of rehabilitation as a continuous and recursive process, applicable in many service settings, which could be independent of the discharge process. He considered the interaction between the individual and the environment to be particularly important. Rehabilitation should entail both working with patients to enhance their confidence and coping skills and the provision of such "prosthetic environments" and social, emotional and material support as may be necessary to maintain their optimal level.

The concept of helping a patient to cope with his/her disability is central to rehabilitation and forms the basis of cognitive-behavioural therapy for schizophrenia²⁵. Patients may deny that they are unwell and this may make engaging them in treatment difficult. It is important to approach their problems from their point of view and work with them to reduce the problems they perceive as important. Education about their illness and medication may be helpful. It is also helpful to discuss exacerbating factors, such as stress and substance misuse. Discussion of early signs of relapse, such as unusual behaviour or prodromal symptoms, may enable patients to prevent a serious relapse. The involvement of family and friends is often helpful. If assessment of risk reveals a substantial risk to the patient or others, then patients should be engaged in the process of reducing the risk.

The Royal College of Psychiatrists report on Rehabilitation emphasized the importance of primary, secondary and tertiary disability²⁶. The primary disabilities are emotional, cognitive, motivational and behavioural dysfunction. Secondary handicaps include loss of self-esteem and confidence, social withdrawal and loss of social roles and networks. Unemployment, homelessness, poverty and stigmatization are the tertiary handicaps. Often patients' main concerns are about the secondary and tertiary problems and it is important to address these to achieve an effective treatment plan.

Deegan²⁷ has described how Mental Health Service users emphasize the importance for them of feeling "in the driver's seat"—empowerment. She uses the word "recovery" rather than "rehabilitation", highlighting the fact that patients recover and professionals rehabilitate. She describes the importance of people becoming experts in their own self-care.

THE PRACTICE OF REHABILITATION

Rehabilitation is based on assessment, development of treatment plans and monitoring. In the UK this forms the basis of the care programme approach. The assessments are based on patients' abilities and difficulties in a wide range of areas. These should include the following.

- *Psychiatric illness*: positive and negative symptoms; insight; compliance with medication; self-medication ability.

- *Physical illness*: specific problems, e.g. diabetes, heart disease; eyesight, hearing and dental problems; preventative medicine, e.g. blood pressure monitoring, breast examination.
- *Education*: literacy and numeracy.
- *Daily living skills*: personal hygiene; laundry skills; budgeting; shopping; cooking; use of public facilities; road safety.
- *Family and social contact*: contact with family; expressed emotion; social networks.
- *Day activities*: work; day hospital; day centre.
- *Financial*: welfare rights, Court of Protection.
- *Accommodation*: patients' wishes; needs based on abilities and difficulties.
- *Risk assessment*: assessment of risk to self and others, including children.

It is often difficult to collect all of the above information and it is important to set a date for an initial review, when the information that has been collected by that time is gathered together. The purpose of the initial review is to appraise the patients' strengths and difficulties and to decide what further information needs to be obtained. Preliminary treatment plans and goals should also be set at this stage. A review of medication, including the use of clozapine for treatment-resistant psychosis²⁸ and the possible use of cognitive-behavioural therapy²⁵ and family interventions²⁹ should be explored.

It is important to have a clear system for reviews, and each patient population or treatment environment requires different systems to be developed, but the essential areas that need to be covered remain the same. Within a district or service it is often helpful to standardize the order of the areas (or headings) in which the assessments are made, so that assessments made in one setting can be easily understood by other teams working in other settings. This leads to improved liaison throughout the service and in particular helps communication between the community and the hospital.

An important part of an initial review is to decide on the order in which problems may be tackled. Sometimes the most important problems are worked on first but on other occasions it may be an idea to start with an easy problem to enable the patient to develop some self-confidence. It is important to set and record practical attainable goals.

Sometimes the review forms or care plan forms that are developed automatically tag the assessment of all strengths and difficulties with treatment plans and goals. This practice can lead to difficulties, as many patients have a large number of problems and in practice only three or four treatment plans can be followed at one time.

The other two vital components of a review are to identify individual staff who have responsibility for components of care and to set a date for a further review when new information on abilities and difficulties can be discussed and the treatment plans and goals evaluated. It is helpful to have patients involved in their treatment programme and many review forms have a space for the patient's signature on them, which assists patient involvement.

RATING SCALES IN REHABILITATION

Although there is no alternative to thorough individual assessments, standardized rating scales can be useful. They are used for the planning of services, the coarse screening of populations to draw up short lists of long-stay patients who may benefit from more intensive assessment of resettlement potential, and for obtaining a quantified assessment of a patient's progress. There are many scales available for assessments in rehabilitation, but few of these have been subjected to investigations of their reliability and validity. The REHAB: Rehabilitation Evaluation³⁰

is an exception to this and is probably the most widely used scale in the UK at present. It is a scale that has been specifically designed for use with long-stay patients and was developed during the course of a 7 year research study. On the basis of behaviour, over 1 week raters complete 23 items in total; seven three-point response scales indicate the frequency of occurrence of difficult or embarrassing behaviours and produce a deviant behaviour score (DV); aspects of social and everyday behaviour are rated in 16 items, each using a visual analogue response format, with written descriptions at each end on which the rater makes a mark that is scored 0–9 using a template. These scores are aggregated to yield the general behaviour score (GB), which can be used to categorize patients as "discharge potential", "moderate handicap" or "severe handicap". There are also five scale scores—social activity, speech disturbance, speech skills, self-care and community skills. The Handbook for the scale is particularly useful, in that it enables one to compare the patient's scores with scores in other populations, so that one can estimate the proportion of people with similar scores in the hospital or the community. Sensitivity to change has also been demonstrated. Provided that raters are properly trained, good reliability levels can be achieved and, once training has been undertaken, the scale itself is brief and easy to complete and score.

CONCLUSION

In conclusion, long-term outcome studies have demonstrated that schizophrenia has a much more heterogeneous outcome than had previously been understood, with many patients making substantial improvements over the years. The effects of environment on outcome are complex and require further long-term evaluation. The aim of rehabilitation is to enable patients to cope with their illness and make the most of their lives, by reducing symptoms and increasing abilities whenever possible and by providing supportive services to match individual need. It is important to see problems from the patients' point of view. In practice this is achieved through a series of reviews which are a focus for assessment, goal planning and evaluation.

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Treatment of Late-onset Psychotic Disorders

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As the world population ages, a greater proportion of the population will be over the age of 65. Psychopathology can present in various ways in the elderly. Psychoses, as defined in DSM-IV, consist of “delusions or prominent hallucinations occurring in the absence of insight into their pathological nature”¹. Hallucinations can be visual, auditory, tactile and/or olfactory in nature. Delusions or falsely-held beliefs are at times very difficult to distinguish from reality. They may require corroboration from caregivers to authenticate their psychotic nature.

Psychotic disorders can present in the elderly as either chronic or acute conditions. The disorders can include early- and late-onset schizophrenia, schizoaffective disorders, delusional disorder, mood disorders with psychotic symptoms, delirium and dementias with psychosis. Furthermore, the etiology of psychosis can be the result of medical conditions, such as Parkinson’s disease or neoplasms, as well as drugs or other substances (Table 95.1).

EPIDEMIOLOGY

The geriatric data from the Epidemiologic Catchment Area study showed, at three sites, a 6 month prevalence for schizophrenia and schizophreniform disorders of 0.2–0.9%; cognitive impairment, including organic psychosis, was 16.8–23%². Reported prevalences of psychotic disorders in a nursing home population was 10% and in a community-based sample in Sweden 4.7%^{3,4}. Patients with Alzheimer’s disease may also exhibit psychotic symptoms, with reported prevalences as high as 63%⁵.

Patients with late-onset schizophrenia (LOS) present with their psychotic symptoms after the age of 45. LOS tend to be predominantly women, and have better premorbid functioning compared to younger schizophrenics⁶. Women with late-onset compared to early-onset schizophrenia tend to have more severe positive symptoms and fewer negative symptoms when compared to men with late onset of the disorder⁷. Women also tend to have less social withdrawal, better premorbid functioning and a gradual decline in functioning^{8,9}. Patients with late-onset schizophrenia tend to have less affect flattening and formal thought disorder⁶. Aging patients who suffer from schizophrenia may have less intensity of their symptoms¹⁰.

Delusional disorders are defined as non-bizarre falsely-held beliefs with minimal hallucinations¹. Delusional disorders often have their onset in mid- or late life¹¹ and tend to affect men earlier than women¹². Low socioeconomic status, immigration, hearing loss and bedfast status are some of the risk factors for developing a delusional disorder^{13–16}. Psychotic symptoms seen in dementias can consist of hallucinations or delusions.

Table 95.1 Psychotic disorders manifested in the elderly

Schizophrenia
Late-onset
Aging patients with early onset
Schizoaffective disorder
Psychosis NOS
Delusional disorder
Dementia with psychotic symptoms
Alzheimer’s dementia
Vascular dementia
Lewy body dementia
Mixed dementias
Delirium
Psychoses due to a general medical condition
Psychoses due to a substance
Mood disorders with psychotic symptoms

NEUROCHEMICAL HYPOTHESIS

The dopamine hypothesis of schizophrenia associates an increase in the activity of dopamine in various cortical areas that are concerned with positive, negative and cognitive symptoms of schizophrenia, as well as side effects seen with the use of neuroleptics. An increase in the level of dopamine activity in the mesolimbic pathways of the brain would be associated with the psychotic symptoms seen, i.e. hallucinations and delusions. Blockade in activity at the mesocortical pathway would be associated with cognitive symptoms and may account for worsening of the negative symptoms seen with conventional antipsychotics. Blockade of dopamine at the tubuloinfundibular tract is associated with increases in prolactin levels, a troublesome side effect which, in younger patients, may lead to non-compliance and discontinuation of treatment.

DIFFERENTIAL DIAGNOSIS

Prior to initiating treatment, it is essential to conduct a biopsychosocial evaluation of the patient. Delirium is often the cause of an acute onset of psychotic symptoms in the elderly and must be foremost in one’s differential assessment. Delirium, defined as an acute mental status change with waxing and waning of the levels of consciousness, often presents with hallucinations, predominantly visual, as well as delusions.

Medical conditions in the elderly that can cause delirium include: infection (frequently urinary tract infections); metabolic disorders (thyroid disease, diabetes); electrolyte imbalances; pain;

Table 95.2 Differential diagnosis of psychotic disorders in the elderly

Medical conditions
Infections
Urinary tract infections
Pneumonia
Viral
Electrolyte imbalance
Endocrine disorders
Cardiovascular events
Myocardial infarctions
Arrhythmias
Neurological
Transient ischemic attacks
Seizures
Strokes
Neoplasms
Other events
Urinary retention
Impaction

acute events, such as myocardial infarctions, strokes, transient ischemic attacks; and exacerbation of chronic conditions, such as chronic obstructive pulmonary disease. Furthermore, medications are frequently part of, if not the underlying etiological cause of, delirium. Medications may contribute as a result of their side-effects profile, toxicity or mechanism of action. Inquiries should be made about the use of other substances, including alcohol or illegal drugs. Corticosteroids, digoxin, anticholinergic drugs and dopaminergic agents are just some of the medications that may be implicated.

The elderly are very sensitive to the side effects of medications, especially to the anticholinergic side effects of drugs. Many medications have anticholinergic side effects, including many antipsychotics, both typical and atypical. The untoward peripheral anticholinergic effects of dry mouth, constipation, urinary retention and dry eyes are troublesome to the elderly and may further aggravate pre-existing physical conditions. Untoward central effects of these medications include the worsening of confusion and cognitive functioning.

TREATMENT OF PSYCHOSIS

One should keep in mind when treating the elderly the old axiom of “do no harm”. The risk:benefit ratio of all treatments prescribed to elderly patients must always be assessed. When choosing an antipsychotic, one should take into account treatment history and susceptibility to potential side effects, as well as family history and familial response to medications. An important principle to apply when initiating treatment is to “start low and go slow”. Treatment should be initiated with doses of half to one-third adult starting dose, remembering that most elderly patients with psychotic symptoms require much lower doses than younger patients.

Age-related bodily changes affect the pharmacokinetics of neuroleptics in the elderly. Absorption of medications may be altered by changes in gastric acidity and emptying and changes in blood flow. Age changes the body’s composition, causing an increase in body fat with an associated decrease in lean body mass and total body water. In addition, there are decreases in liver mass and blood flow and changes in renal blood flow and function. All these changes affect the absorption, metabolism, distribution and clearance of neuroleptics.

The phenothiazides are lipophilic substances, well absorbed, with an extensive first-pass metabolism in the liver. In the elderly, as a result of the increase in total body fat, there is an increase in the volume of distribution of drugs. This leads to an increased half-life for those substances that are lipophilic. Thus, the elderly should be given lower doses of lipophilic drugs.

Neuroleptics are the treatment of choice for psychosis. They have recently been divided into typical (conventional) and atypical antipsychotics, based on their capacity to treat the positive and negative symptoms of schizophrenia as well as their potential to cause neurological side effects. All antipsychotics have the same level of efficacy. It may be relevant to recall that 100 mg chlorpromazine is equivalent to 1 mg haloperidol and 0.5 mg risperidone.

Typical or “conventional” antipsychotics can be divided into groups, based on their potency—high, intermediate and low potency. High-potency antipsychotics include haloperidol, fluphenazine and loxapine. High-potency neuroleptics have greater affinity for the dopamine receptors and less affinity for the muscarinic and α -receptors. High-potency neuroleptics are more likely to be associated with a higher incidence of extrapyramidal symptoms, akathisia, acute dyskinesia and parkinsonism, thus limiting their use in the elderly. Intermediate potency neuroleptics include perphenazine, loxapine and molindone. The atypical neuroleptics are clozapine, risperidone, olanzapine, quetiapine and ziprasidone. Low-potency antipsychotics, such as thioridazine and chlorpromazine, have a higher affinity for muscarinic, histaminic and α -adrenergic receptors and furthermore are more likely to produce increased sedation, orthostatic hypotension and anticholinergic side effects. They should be prescribed with caution in the elderly.

Limitations of Typical Neuroleptics in Geriatrics

Typical antipsychotics block various receptors that have the potential to cause side effects which can limit their use in the elderly. Dopaminergic blockade is associated with acute and long-term neurological side effects. Acute neurological side effects are extrapyramidal side effects (EPS), which include parkinsonism (resting tremors, rigidity, bradykinesia and gait disturbances), akathisia and dystonias and long-term effects of tardive dyskinesia, as well as the potential for neuroleptic malignant syndrome (NMS). Histaminergic blockade is associated with sedation and weight gain. Their quinidine-like cardiac effects are associated with the potential for arrhythmias. α -Adrenergic blockade leads to orthostatic hypotension. In addition, the muscarinic blockade leads to anticholinergic side effects.

Neuroleptic induced parkinsonism (NIP) is a potential concern with the use of neuroleptics in the elderly. The reported prevalence of NIP in patients aged over 60 on neuroleptics is 50%¹⁷. One study that looked at the use of low-dose neuroleptics and the incidence of NIP in the elderly found that 32% of patients developed NIP on an average daily dose of 43 mg chlorpromazine¹⁷. In addition, the risk factors contributing to the incidence of NIP were older age, instrumental tremor at baseline, EPS, the type of neuroleptic administered and the severity of dementia^{17,18}. Parkinsonism can increase the risk of dependency, falls and fractures¹⁹. These often troublesome side effects can be treated either by reducing the neuroleptic dose, switching to an atypical agent, or using anticholinergics such as benzotropine or trihexyphenidyl. Care needs to be taken when using antiparkinsonian medications in the elderly, who are very sensitive to the anticholinergic side effects.

Akathisia is another neuroleptic side effect that can be difficult to address. Akathisia is characterized by increased restlessness and psychomotor activity, with an inability to sit still. Akathisia is

Table 95.3 Medications/substances associated with delirium

Corticosteroids
Digoxin
Pain medications
Opioids
Muscle relaxants
H2 blockers
Anticholinergics
Benzodiazepine withdrawal
Alcohol withdrawal
Alcohol intoxication

often mistaken for worsening of the psychotic symptoms, leading one to erroneously increase the neuroleptic dose, when actually lowering the dose may address this troublesome side effect. In addition, akathisia can be treated with benzodiazepines and/or β -blockers.

Tardive dyskinesia (TD) is a potentially irreversible abnormal involuntary choreiform movement disorder. In the elderly, the 3 year cumulative incidence of severe TDs was found to be 2.5% after 1 year, 12.1% after 2 years and 22.9% after 3 years²⁰. Another study reported the cumulative rates for TD to be 25% after 1 year, 35% after 2 years and 53% after 3 years²¹. Factors that were predictive of TD included higher daily dose at study entry, greater cumulative amounts of prescribed neuroleptics, greater severity of worsening negative symptoms, and the presence of early EPS^{21,22}. Caution should be used when administering conventional neuroleptics in the elderly and they should only be prescribed when necessary and at the minimal effective dose²⁰.

Neuroleptic malignant syndrome (NMS) is a serious side effect with the potential to be lethal. NMS presents with symptoms of muscle rigidity, fever, autonomic instability, fluctuating levels of consciousness and elevations in CPK and white blood cell counts. NMS can be seen with the use of all neuroleptics, including the atypical²³.

The potential cardiovascular effects of these drugs include orthostatic hypotension and QRS prolongation, leading to an increased risk for torsade de pointe. Orthostasis is of especial concern in the elderly who, due to physiological changes, are already at increased risk. In addition, in those patients who are already suffering from orthostasis or who are being prescribed medications known to have pressure-lowering effects, low-potency neuroleptics should be avoided. Orthostatic hypotension places the elderly at increased risk of falls, leading to increased morbidity and mortality.

The anticholinergic effects of these drugs can have both troubling and undesired peripheral and central effects. Peripheral side effects include dry mouth and eyes, blurred vision, constipation, and urinary retention, which is of especial concern in males who have prostatic hypertrophy. Central effects include worsening of cognition, confusion and/or delirium^{3,24}. One should be aware

that the atypical antipsychotics clozapine and olanzapine also have anticholinergic side effects.

An additional side effect of concern with the use of antipsychotics is photosensitivity in those patients who are on chlorpromazine. Patients may experience irreversible degenerative pigmentation retinopathy, caused by doses of thioridazine greater than 800 mg/day. Weight gain is of especial concern in those patients with an obesity problem or who are known to gain weight easily on medications. Weight gain can be seen with the use of both conventional and atypical antipsychotics.

Atypical Antipsychotics

Atypical antipsychotics are so defined because they are effective at treating both the positive and the negative symptoms of schizophrenia, while having a lower incidence of extrapyramidal symptoms and tardive dyskinesia. FDA-approved atypical antipsychotics are clozapine, risperidone, olanzapine, quetiapine and ziprasidone.

Clozapine

Clozapine was the first atypical antipsychotic approved for use in the USA. Compared to typical neuroleptics, it has a higher affinity for the dopamine D2 receptor, being more selective for the mesolimbic and mesocortical pathways. Clozapine continues to have a favorable neurological side-effect profile but is reported to have the following untoward effects: drowsiness, sedation, hypersalivation, tachycardia, dizziness, constipation, nausea and vomiting, the most concerning being agranulocytosis and seizures. It is recommended that clozapine treatment should be initiated only after a patient has failed two trials with conventional antipsychotics. In a double placebo-controlled study of six patients, clozapine was found to have similar efficacy as in younger patients. Although it was felt to be fairly safe in the elderly, the most frequent side effects were sedation, confusion and agranulocytosis²⁵.

Concern exists that the elderly may be at greater risk for agranulocytosis. In an Australian elderly study, the occurrence of agranulocytosis was found to be 4% compared to 0.25% in younger patients²⁵. Increased sedation, hypersalivation, bradycardia, postural hypotension and delirium are frequent side effects reported in the elderly with the use of clozapine^{26–28}. Another potential side effect is weight gain. One author found that 75% of the patients prescribed clozapine had gained an average of 7.5 kg²⁹. The potential for agranulocytosis requires constant monitoring of blood, which may be difficult and costly in the elderly. Clozapine was tolerated as well as chlorpromazine in one study. In one open label report in 300 older adults, the discontinuation rate for clozapine was 43% and agranulocytosis occurred in two non-fatal cases³⁰.

Clozapine should be considered once a patient has failed two trials of conventional neuroleptics. When prescribing clozapine, it is prudent to initiate treatment with a dose of 6.25 mg/day, followed by weekly titration of 6.25 mg/day until a therapeutic effect is achieved and/or side effects develop^{13,31}. Daily doses may range from 6.25 to 400 mg³¹. Clozapine should be initiated slowly and titrated slowly. Chengappa²⁶ has reported that the rapid titration of this drug can lead to drug intolerance and poor response.

Table 95.4 Recommended doses in the elderly

Drug	Initial dose	Maximum dose
Clozapine	6.25 mg/day	50–100 mg/day
Risperidone	0.25–0.50 mg/day	2 mg/day
Olanzapine	2.5 mg/day	5–10 mg/day
Quetiapine	25 mg/day	100–150 mg/day
Ziprasidone	20 mg/day	80–160 mg/day

Risperidone

Risperidone, a serotonin–dopamine antagonist, is an atypical antipsychotic that is a benzisoxazole derivative³². Risperidone has been found to be effective for the treatment of both positive and negative symptoms of schizophrenia, with minimal EPS. Risperidone has been studied extensively in the elderly^{33,34}. It is a potent antagonist of the 5-HT receptor and less so of D2, with α -receptor blocking. In addition, it has minimal histaminergic blockade and practically no affinity for cholinergic receptors³⁵.

Side effects seen with this drug include orthostatic hypotension, sedation, fatigue and palpitations. Risperidone has dose-dependent incidence of EPS and TD. The incidence of sedation is in the range 6–15%^{36–38}. There are no cardiac or laboratory abnormalities associated with this drug. Hypotension can be seen and should be carefully monitored in patients with pre-existing hypotension or who are on medications that have this effect. The 1 year incidence of TD seen in a large sample of elderly patients treated with risperidone was 2.6%³⁹.

Risperidone is the atypical antipsychotic most widely studied in the elderly. Most studies have favored the use of lower doses, finding them to be efficacious. In a study of patients who were diagnosed with dementia with behavioral disturbances, risperidone was found to be effective at doses of 1 mg/day; more adverse events were seen at doses of 2 mg/day³⁴. These authors recommended the use of a serotonin–dopamine agonist when treating elderly patients with psychotic symptoms³⁴. In a population of American veterans the average dose of risperidone was 3.6 mg/day compared to an average olanzapine dose of 10.2 mg/day⁴⁰ that EPS were reduced from baseline.

Risperidone has been found to be effective in treating both the negative and positive symptoms in patients with the diagnosis of schizophrenia and schizoaffective disorders. In a prospective open-label, 12 week study of 103 elderly patients who were diagnosed with schizophrenia or schizoaffective disorders the mean dose was 2.4 mg/day. An overall improvement of 45% was seen for both PANNS and CG³⁷. The most frequently reported adverse events were dizziness, insomnia, agitation, somnolence, constipation and EPS. The authors concluded that risperidone was safe in this population but that doses should not be lower than 3 mg/day. Various authors have found risperidone to be a safe and efficacious treatment in this population^{33,37,38}. In the medically ill elderly with psychotic symptoms, risperidone is also an effective and safe treatment. It does demonstrate dose-dependent EPS^{37,41}. Risperidone treatment should be initiated at low doses (0.25 mg/day) and slowly titrated to a target dose of 1.5 mg/day for patients with dementia with psychotic symptoms. In elderly patients with psychosis, the optimal dose should be 2 mg/day. Risperidone in the elderly is an effective safe first-line treatment of psychosis.

Olanzapine

Olanzapine is a thienobenzodiazepine analog with an *in vitro* receptor affinity profile similar to clozapine. It is considered to be an atypical antipsychotic, as it is more effective than haloperidol in the treatment of both positive and negative symptoms of schizophrenia with minimal EPS⁴². Olanzapine affects the dopamine D1 and D4 receptors and the serotonin 2a and 2c receptors; it is well absorbed, with a mean half-life of 30 h, and is metabolized by the cyp 1a2, cyp2 D6 and flavin mono-oxygenase system. The CYP2D6 is a minimally involved system, thus there should be minimal concern for drug interactions with potent inhibitors of this system. Olanzapine has no active metabolite⁴².

The most frequent untoward events seen with olanzapine are hypotension, constipation, weight gain, somnolence, agitation

and dizziness⁴². It does have anticholinergic side effects, and can cause significant weight gain, which may be an advantage for those elderly patients who have suffered significant weight loss. The optimally recommended dose range is 2.5–15 mg/day.

An ad hoc study reviewing the olanzapine data in those over the age of 65 with psychotic disorders found no significant difference between the elderly and younger patients. Dose comparisons were similar for patients younger and older than 65^{43,44}. A study using olanzapine in patients with dementia of the Alzheimer's type showed no difference in efficacy from the placebo group at doses in the range 1–8 mg/day. This same study found no difference in liver function test, EPS, leukopenia or orthostasis when compared to placebo⁴⁵. Although olanzapine appears to have anticholinergic side effects, these were not significantly different from those seen in comparison to a placebo group⁴³. Olanzapine was found to be well tolerated and safe in the elderly. Doses used are similar to those for younger patients; initial doses should be 2.5–5.0 mg/day. Olanzapine has the added advantage of once daily dosing.

Quetiapine

Another novel antipsychotic recently FDA-approved for use in the USA is quetiapine. Quetiapine is a dibenzothiazepine derivative. It exhibits higher affinity for 5-HT₂ receptors than for dopamine D2 receptors and is reported to have greater affinity for the mesolimbic than the nigrostriatal sites, thus accounting for its effect on positive symptoms with minimal EPS^{46,47}. Quetiapine is reported to treat both the positive and negative symptoms of schizophrenia^{46,47}. The incidence of EPS has been found to be minimally no different than placebo.

In a study of elderly patients with psychotic disorders, the average dose of quetiapine was 100 mg/day. In that same study the most frequent side effects found were somnolence (32%), dizziness (14%), postural hypotension (13%) and agitation (11%). However, from a cardiovascular perspective, the patients did exhibit a slight increase in their heart rate but there were no changes in the QRS complex. No hematological changes were noted, except for a non-significant slight increase in t4 not associated with substantial changes in the mean thyroid-stimulating hormone levels. Quetiapine was rarely associated with weight gain. These authors recommended starting at a dose of 25 mg/day and titrating to a target dose of 100 mg/day in divided doses. Most patients will be treated with doses in the range 100–300 mg/day, with an occasional patient requiring higher doses^{46,48}.

Ziprasidone

A new atypical antipsychotic currently awaiting FDA approval in the USA is ziprasidone. Ziprasidone is reported to have affinity for the dopamine D2 and D3 receptors and the agonist of 5-HT₂. It has negligible affinity for the other receptors of concern in the elderly, including α -2 and β -muscarinic; it moderately inhibits the reuptake of serotonin and norepinephrine but weakly effects the reuptake of dopamine^{49,50}. This drug is potentially effective for both positive and negative symptoms and depression, with a low likelihood of causing EPS⁴⁹.

The following studies have looked at the use of ziprasidone in younger patients, the oldest being 64. Patients with acute exacerbation of schizophrenia and schizoaffective disorder had a better response than placebo on both 80 mg/day and 160 mg/day⁵⁰. Often patients with these disorders are suffering from depression; in this same study 160 mg of ziprasidone was found to be effective to treat depression⁵⁰. Other studies have reported that a dose of 120 mg/day was more effective than 40 mg/day. The side

effects reported included dyspepsia, transient somnolence, constipation, nausea and abdominal pain⁴⁹. Ziprasidone was associated with a lower incidence of EPS and postural hypotension, with no weight gain or laboratory abnormalities⁴⁹.

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Risk Factors for Dyskinesia in the Elderly

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TARDIVE DYSKINESIA

Clinical Features

In the mid-1950s, within a few years of the introduction of antipsychotic drugs, clinicians drew attention to “neurotoxic reactions” to this treatment. In addition to parkinsonian features, involuntary orofacial movements were noted^{1,2}. Sigwald *et al.*³ are usually credited with the first detailed description of “facio-bucco-linguomasticatory dyskinesia” associated with these drugs. In the early 1960s, Faurbye *et al.*⁴ coined the term “tardive dyskinesia” for this condition.

Abnormal involuntary orofacial movements remain the most familiar and prevalent features of tardive dyskinesia. The movements are irregular, stereotyped and choreic in nature, and tend to involve the tongue, lips, jaw and face, including the peri-orbital areas. The movements observed include protrusion or twisting of the tongue, lip smacking, cheek puffing, pursing and sucking actions of the lips, and chewing and lateral jaw motions. The particular combinations of movement seen vary considerably between patients but tend to be relatively consistent for each individual.

In addition to these orofacial phenomena, most descriptions of tardive dyskinesia include a variety of trunk and limb movements. These are typically choreiform or choreoathetoid in type, although athetosis of extremities, and axial and limb dystonia, are often listed as part of the syndrome, as are abnormalities of gait and trunk posture, such as lordosis, rocking and swaying, shoulder shrugging and rotary movements of the pelvis. Grunting and respiratory arrhythmias are also seen.

The notion that orofacial and limb and trunk dyskinesia represent distinct pathophysiological entities with different risk factors and clinical correlates⁵ seems to hold true in older patients⁶. For example, Paulsen *et al.*⁷ studied middle-aged and elderly outpatients starting antipsychotic drug treatment and obtained systematic follow-up data over 2 years. They found that the cumulative incidences and the significant predictors identified differed for the two subsyndromes.

Prevalence, Incidence and Natural History

While tardive dyskinesia is a common problem in those individuals receiving antipsychotic drug treatment long-term, the majority will not exhibit the condition. The reported prevalence figures vary widely, from 0.5% to 56%^{8,9}, reflecting variables such as the age of the sample studied (see below) and the sensitivity of the rating instrument used. Overall, the literature suggests that

only 20% or so of drug treated patients will develop the problem, and of these, the dyskinesia is likely to be serious in less than 10%. Gardos *et al.*¹⁰ concluded that the severe tardive dyskinesia is very uncommon, occurring in approximately 1/100–1/1000 patients with the condition. Nevertheless, in elderly psychiatric inpatients, prevalence figures of around 50% or greater are not uncommonly reported⁶.

The prospective study of Kane *et al.*^{11,12} suggested that new cases of tardive dyskinesia can occur at a rate of 3–4% a year in the first few years after starting antipsychotic medication. However, there is only limited information available to guide the clinician as to who might be most at risk. Tardive dyskinesia was originally considered irreversible, although follow-up studies^{13,14} have revealed that it is not a progressive disorder, but rather tends to fluctuate in severity over time. Spontaneous remissions are relatively common, particularly in younger patients. Such a view of the natural history of tardive dyskinesia in patients on chronic drug treatment was reinforced by the results of two 10 year follow-up studies by Yagi and Itoh¹⁵ and Casey and Gardos¹⁶.

To some degree, these findings might be seen as relatively reassuring, allaying earlier fears that tardive dyskinesia was a major iatrogenic condition that would constrain the use of long-term antipsychotic drug therapy. However, while the condition itself is rarely disabling, it can be socially stigmatizing. Further, severe tardive dyskinesia can be a troublesome problem, particularly in the elderly. Orofacial dyskinesia can interfere with eating and swallowing, render speech unintelligible, and cause breathing difficulties, sometimes leading to dysphagia or choking^{17–19}. Trunk and limb dyskinesia may be associated with disturbances of gait that can lead to falls and injury²⁰.

RISK FACTORS

Advancing Age

In numerous studies, advancing age has been clearly shown to be a major vulnerability factor for tardive dyskinesia. It is associated not only with an increased occurrence of tardive dyskinesia but also with greater severity and a reduced likelihood of spontaneous remission^{9,21}. Studies involving geriatric patients have consistently found a higher incidence of tardive dyskinesia with conventional antipsychotics, even at low doses^{22,23}. Examining pooled epidemiological data has revealed a strong linear correlation between age and both the prevalence and severity of tardive dyskinesia²⁴. However, duration of exposure to antipsychotic drugs is a confounding variable, as older patients are likely to have received

drug treatment for longer; i.e. age-group and length of time on drug treatment would be expected to be linked. Nevertheless, clinical studies have consistently shown an effect of age independent of that attributed to length of drug treatment^{9,13}. Further, it would seem that the older patients are when they first receive antipsychotic medication, the more likely they are to develop the condition early in treatment²⁵. This was dramatically demonstrated in a prospective study of tardive dyskinesia in patients 55 years or older who were just beginning antipsychotic drug treatment²⁶. These investigators initially presented preliminary data on 160 patients²⁷ and subsequently reported on the complete study group of 261 patients²⁶. Patients were assessed at 3 month intervals. After approximately 1, 2 and 3 years of cumulative exposure to antipsychotic medication, the incidence figures for tardive dyskinesia in the larger sample were 25%, 34% and 53%, respectively. These investigators concluded that the rate of tardive dyskinesia for patients over 50 years of age starting antipsychotic drug treatment was three to five times that reported in younger adults.

Much of the variation in incidence and prevalence figures reported in the published literature is likely to be due to age differences in the sample studied⁹. The results of a 3 year follow-up study of tardive dyskinesia by Barnes *et al.*¹³ suggested that patients receiving antipsychotic drugs may be most likely to develop the condition in their sixth decade. Jeste *et al.*²⁸ calculated that in the sample of chronically ill, schizophrenic inpatients they studied, the probability of having tardive dyskinesia was greater than 0.4 in those patients over 65 years old. In their review of the literature, Smith and Baldessarini²⁴ pointed out that the rise in prevalence with age may partly reflect the relative persistence of tardive dyskinesia in older patients. They concluded that tardive dyskinesia in those less than 60 years of age was over three times more likely to remit spontaneously. However, as the relevant studies included patients either withdrawn from antipsychotic medication or on extended "drug holidays", this conclusion partly reflects the reversibility of the condition after drugs are discontinued. Younger patients appear to possess a greater recovery potential when medication is withdrawn, but whether they show a higher spontaneous remission rate than older patients when maintained on chronic antipsychotic drug therapy remains unconfirmed. Work with an animal model of tardive dyskinesia suggests that spontaneously remitting dyskinesia emerging early in drug treatment can become irreversible with prolonged drug therapy²⁹.

Antipsychotic Medication

While it seems plausible that the relationship between exposure to antipsychotic treatment and vulnerability to tardive dyskinesia varies with age, such a relationship has not been systematically tested. In the minority of studies where a relationship between specific drug variables and tardive dyskinesia has been found, it has tended to be within the first few years of treatment^{9,11}, particularly in studies of elderly populations³⁰⁻³². For example, Jeste *et al.*²¹ studied 266 patients over 45 years of age receiving conventional antipsychotic medication. At baseline, the median duration of total lifetime antipsychotic exposure was only 21 days. Over the first 3 years, the cumulative incidence of tardive dyskinesia was 26%, 52% and 60%, respectively. The main risk factors for the appearance of tardive dyskinesia included baseline duration of antipsychotic treatment of over 90 days, and the cumulative amount of antipsychotic medication prescribed, particularly "high-potency" drugs. This group of investigators³³ also examined risk factors specifically for severe tardive dyskinesia. In a prospective study of 378 older neuropsychiatric patients over 3 years, the incidence of severe tardive dyskinesia was 2.5%

in the first year, 12% in the second and 23% in the third. As in the earlier study, higher daily doses of antipsychotic medication at baseline and greater cumulative amount of prescribed antipsychotic emerged as risk factors. In the study by Woerner *et al.*²⁶ mentioned above, higher mean daily dose of antipsychotic and cumulative dosage were also found to be associated with a greater risk of developing tardive dyskinesia.

The mechanisms whereby advancing age exerts an influence on susceptibility to tardive dyskinesia remain speculative. It is possible that neuropathological and/or neurochemical changes occurring with age may be associated with an increasing prevalence of tardive dyskinesia. Neuronal damage or degeneration, receptor changes and the reduced efficiency of adaptive homeostatic processes may well underlie the vulnerability of certain drug-induced neurological side effects in older patients. The development of supersensitive post-synaptic dopamine receptors in the basal ganglia remains the accepted explanation of the pathophysiology of tardive dyskinesia. However, such supersensitivity would seem to be an inevitable consequence of prolonged antipsychotic drug treatment, and is therefore insufficient to explain why only a proportion of patients on long-term medication develop dyskinesia³⁴. An interaction between drug-induced receptor changes in the striatum, and age-related degenerative effects in the nigrostriatal system may be necessary for the appearance of tardive dyskinesia.

Pharmacokinetic mechanisms could also be relevant. Age-related changes in the absorption, metabolism and excretion of drugs may lead to higher plasma drug levels and delayed clearance. Plasma levels of antipsychotic drugs have been found to be raised in the elderly, compared with younger patients^{35,36}.

Atypical Antipsychotics

The newer, atypical antipsychotics are characterized by a lower liability for extrapyramidal side effects and, for some of these agents, evidence is emerging for a lower risk of tardive dyskinesia³⁷. However, thus far, there have only been a few studies addressing the safety and tolerability of these drugs, at appropriate dosage, in the elderly. For reasons addressed briefly above, the findings of clinical trials in young adults may not be reliably extrapolated to the elderly, for whom dosage requirements tend to be lower^{38,39}. Preliminary clinical data suggest that newer agents such as risperidone and olanzapine may be relatively well tolerated in older people⁴⁰ and there have been similar claims for low-dose clozapine⁴¹, but well-designed, controlled studies in the elderly are required^{42,43}.

In respect of tardive dyskinesia risk in the elderly, risperidone has been the atypical antipsychotic most commonly studied. Over a period of 9 months, Jeste *et al.*⁴⁴ found a significantly lower incidence of tardive dyskinesia in older patients treated with risperidone as opposed to haloperidol. The patients were middle-aged or elderly (mean age 66 years) and had not received previous treatment with antipsychotics. "Clinically comparable" groups receiving treatment with either risperidone or haloperidol were compared. The median dose for both drugs was 1mg/day. The haloperidol-treated patients proved to be significantly more likely to develop tardive dyskinesia. A further study by the same group of investigators⁴⁵ addressed the risk of persistent tardive dyskinesia with risperidone in elderly patients with dementia. In an open study, a sample of 330 patients (mean age 82.5 years) received risperidone in flexible dosage (mean modal dose 0.96 mg/day) over a year. Amongst the 255 patients with no evidence of tardive dyskinesia at baseline, the 1 year cumulative incidence of the condition was 2.6%, and patients who had exhibited dyskinesia at baseline experienced significant improvement. The investigators concluded that such

an incidence was much lower than would be expected with conventional antipsychotic treatment in such patients. A similar conclusion was reached by Davidson *et al.*⁴⁶, who conducted an open study of risperidone in a sample of 180 elderly, chronically-ill, psychotic patients (median age 72 years) over 1 year; 97 of those in the sample received risperidone for the full 12 months. At the endpoint, the mean dose of risperidone was 3.7 mg/day. Only six cases of persistent tardive dyskinesia emerged during the study period, representing an incidence of 4.3%.

Preliminary evidence also raises expectations of a relatively lower risk of tardive dyskinesia in the elderly with olanzapine^{18,47} and possibly quetiapine¹⁸.

Gender

A relatively consistent finding has been that women show a greater prevalence of severe dyskinesia, although there is evidence to suggest that this is limited to the geriatric age range, that is, over 60 years of age^{9,48,49} and the more severe cases^{50,51}. The ability to detect a significant effect for gender is probably related to the size of the study sample and the base rate of tardive dyskinesia in the population being studied. Nevertheless, Smith and Dunn⁵² identified 13 studies reporting statistically significant differences supporting female vulnerability, with the average unweighted female:male prevalence ratio in the patient samples being 1.69.

For example, Ramsay and Millard⁵³, in a sample of 426 elderly subjects, predominantly geriatric or psychogeriatric inpatients, found dyskinetic movements in 12.5% of women but only 7.6% of men. In a sample of patients over 65 years with senile dementia of the Alzheimer type, O'Keane and Dinan⁵⁴ found that significantly more females (89%) had evidence of orofacial dyskinesia compared with the males (60%). However, the prospective study of tardive dyskinesia in the elderly, conducted by Woerner *et al.*²⁶, failed to find a significant relationship between sex and incidence of the condition. Indeed, the incidence was slightly higher for males, although they were significantly younger than the females.

Richardson *et al.*⁵⁵ reported that females showed an increase in prevalence of tardive dyskinesia in all age groups up to and beyond 75 years, while males only demonstrated an increase up to 75 years, with a decline subsequently. The differences in prevalence between the sexes was minimal up to 64 years using mild severity to define a case, but female prevalence was substantially higher when a criterion of moderate to high severity was applied. These findings support the view that the magnitude of sex differences in tardive dyskinesia is dependent on the severity of the criteria used to define the disorder, as well as the age of the sample^{50,51}.

As a vulnerability factor, gender is far weaker than age, with which there seems to be an interaction. There is no clear explanation for the female preponderance, although it has been suggested that neuroendocrine factors may be relevant. Oestrogen and prolactin may influence striatal dopamine function⁵⁶, and the reduced ovarian function in post-menopausal women, with low oestradiol levels, may increase vulnerability to tardive dyskinesia^{57,58}. However, rather than a direct reflection of gender, it has been variously speculated that female sex is a proxy variable for longer duration of hospitalization, higher drug dose and longer duration of treatment.

Organicity

Reviews of the risk factors for tardive dyskinesia⁵⁹⁻⁶¹ suggest that in patients with schizophrenia, evidence of "organicity" is a

marker of vulnerability to tardive dyskinesia. Putative, indirect indices of organicity, such as soft neurological signs and cognitive impairment, computed tomography (CT) evidence of structural brain pathology and the presence of negative symptoms, have all been found more commonly in patients with tardive dyskinesia than those without. Thus, ageing may be a predisposing factor for tardive dyskinesia only insofar as it increases the likelihood of organic brain dysfunction.

Hunter *et al.*⁶² and Crane and Paulson⁶³ surveyed mixed psychiatric inpatient populations and found tardive dyskinesia to be more prevalent in patients with organic mental syndromes than in those with schizophrenia. Yassa *et al.*⁶⁴ conducted a similar study, assessing over 300 psychiatric patients treated with antipsychotic drugs. They also found that patients with organic mental syndromes, including those diagnosed as epilepsy with psychosis and mental retardation with psychosis or alcoholism, showed a significantly higher prevalence of tardive dyskinesia compared to those with schizophrenia. A number of organic mental syndromes and neuromedical conditions, such as multi-infarct dementia, strokes and cerebral tumour, are more common in older individuals, and this may partly explain why the prevalence of tardive dyskinesia is greater in elderly samples^{65,66}. However, any such hypothesis should take account of the influence of the range of other psychiatric diagnosis in such samples. For example, Yassa *et al.*⁶⁴ found that patients with a primary bipolar affective disorder had the same prevalence of tardive dyskinesia as those with organic mental syndromes. In their prospective study in the elderly, Woerner *et al.*²⁶ reported that the risk of tardive dyskinesia in individuals with multi-infarct dementia was similar to that for people with mood disorder, while patients with Alzheimer's disease and other organic mental syndromes showed a lower rate.

Psychiatric Diagnosis

Affective Disorder

Recent work has highlighted affective disturbance as a possible relevant variable. Both a positive family history of affective disorder and the presence of depressive features in patients with schizophrenia have been mooted as markers of an increased likelihood of developing both parkinsonian side effects and tardive dyskinesia, but the evidence is tentative at present. However, there is accumulating evidence that patients with primary affective disorder may be at particular risk of tardive dyskinesia if administered antipsychotic drugs long-term. A report by Davis *et al.*⁶⁷ noted the relatively high prevalence of tardive dyskinesia among such patients and this has been confirmed by later studies⁶⁸⁻⁷⁰.

The preliminary data from the prospective study of tardive dyskinesia by Kane *et al.*¹⁹ provides additional support for a diagnosis of affective disorder as a vulnerability factor. A life-table analysis based on the length of drug administration, and comparing the cumulative incidence of tardive dyskinesia in patients with affective or schizoaffective disorder with that of inpatients with a diagnosis of schizophrenia, revealed that the former have a significantly greater incidence after 6 years of exposure to antipsychotic drugs. The incidence figures were 26% for the affective and schizoaffective patients compared with 18% for those with schizophrenia. Further evidence was provided by Mukherjee *et al.*⁷¹, who found persistent tardive dyskinesia in 35% of a sample of bipolar patients who had received maintenance antipsychotic drugs, while no patients without such a drug history had persistent dyskinesia.

Schizophrenia

The schizophrenic illness itself may be a risk factor^{59,61}. Long before the advent of antipsychotic drugs, a variety of motor disorders were described in psychiatric patients, particularly those with catatonic schizophrenia⁷²⁻⁷⁴ and other types of schizophrenia⁷⁵⁻⁷⁷. These movements would seem to be principally disturbances of voluntary motor activity and may be classified as stereotypes and mannerisms, preservative movements, tics, grimaces and general clumsiness, awkwardness and lack of coordination. However, Kraepelin⁷⁸, and then later Farran-Ridge⁷⁵, observed spasmodic movements, mainly involving the orofacial muscles, which they considered choreiform in nature.

Nevertheless, Marsden *et al.*⁷⁹ concluded that true chorea and athetosis were scarce in chronic psychiatric patients before drug treatment, and that much of the motor disorder seen was attributable to organic neurological disorder, such as encephalitis and syphilis. Further, the terminology used is confused, reflecting various conceptual notions of the aetiology of the motor phenomena observed^{80,81}. Kleist⁷³ and Farran-Ridge⁷⁵ commented that the similarity between the manifestations of dementia praecox and epidemic encephalitis was such that difficulties in differential diagnosis could arise. However, there is no doubt that some of the movements described would be scored on current rating scales for tardive dyskinesia if seen now in patients receiving antipsychotic drugs.

There would seem to be three possible interpretations of these observations⁵⁹. First, the type of motor disturbance historically described is not specifically associated with schizophrenia, but rather the product of organic brain disease. The two conditions also occur together when the brain disease is also responsible for symptomatic schizophrenia, as, for example, when schizophrenia appears in patients with encephalitis, Wilson's disease or Huntington's disease, among other conditions⁸². Second, the association of motor disturbance and schizophrenia may be more specific, in that an underlying neuropathological process is capable of producing both psychological and motor impairments. Third, as Kraepelin and Bleuler tended to suggest, the movements may be secondary to the schizophrenic disturbance of will, thought and emotion. However, the distinctions between the three explanations cannot be too sharply drawn, and more than one may be relevant.

In a relevant contemporary study, Owens *et al.*⁸³ compared chronic schizophrenia inpatients with and without a history of antipsychotic drug treatment, and found a similar prevalence of abnormal involuntary movements in the two groups. This finding confirmed the occurrence of spontaneous movement disorder in schizophrenia. However, the contribution of drug therapy was acknowledged, in that when the age difference between the two patient samples was taken into account in further analysis of the data, there was a significant linear relationship between the prevalence of abnormal involuntary movements and exposure to antipsychotic drugs⁸⁴. Grouping movements into clinically recognizable syndromes revealed a particular susceptibility to orofacial dyskinesia in the drug-treated patients. Nevertheless, the schizophrenic illness, at least in some forms, may be seen as a psychomotor disorder with an inherent, increased risk of dyskinesia. Antipsychotic drug treatment may interact with the disease process and age-related cerebral deterioration to hasten or provoke the appearance of such movement disorder^{59,85}.

Dementia

Accepting that organicity is a risk factor for tardive dyskinesia, it might be expected that the neurodegenerative changes of Alzheimer's disease would be associated with a relatively high

risk of spontaneous dyskinesia and also a greater risk of tardive dyskinesia when antipsychotic drugs are administered. Molsa *et al.*⁸⁶ assessed abnormal involuntary movements in 177 patients with Alzheimer's disease, with a mean age of 75 years. In the 143 patients who had never received antipsychotic drugs, the prevalence of dyskinesia was 17%, while for the 34 patients treated with antipsychotic drugs the figure was 53%. As part of a larger study, Ramsay and Milard⁵³ looked at 40 patients on long-stay psychogeriatric wards. Of the 13 (32.5%) patients with dyskinesia, 12 had a history of treatment with antipsychotic drugs. These findings suggest a vulnerability to tardive dyskinesia in dementia patients receiving antipsychotic drugs.

O'Keane and Dinan⁵⁴ assessed 78 patients with an age range of 65-91 years, all of whom fulfilled DSM-III criteria for senile dementia of the Alzheimer type. The scales used included the Mini-Mental State (MMS)⁸⁷ and Abnormal Involuntary Movements Scale (AIMS)⁸⁸. They reported that 62 patients (69%) had evidence of orofacial dyskinesia, a figure the authors noted to be over 10 times that reported in healthy elderly adults, and also considerably higher than the prevalence found in populations of patients with chronic schizophrenia. Orofacial dyskinesia was by far the most common abnormal movement rated. The mean doses of antipsychotic drug for those with and without abnormal movements were modest, and although the former group was receiving a higher mean dosage, the differences between the two groups was not statistically significant.

Molsa *et al.*⁸⁶ found that the severity of orofacial dyskinesia in their sample of patients with Alzheimer's disease increased with the degree of cognitive deficit. Bakchine *et al.*⁸⁹ studied a group of 91 patients with dementia of the Alzheimer type and examined the relationships between primitive reflexes, extrapyramidal symptoms and severity of cognitive impairment. They failed to find any significant relationship between "buccolinguofacial dyskinesias" and a low score on intellectual functioning, but their methodology was criticized by O'Keane and Dinan⁵⁴, who pointed out that they only rated abnormal movement as either present or absent, rather than qualifying the movements using a standardized scale. In their own study O'Keane and Dinan found that there were no significant differences in terms of age or length of illness between those patients with and without abnormal movements, and both groups showed evidence of severe intellectual impairment. However, those patients with abnormal movements had a significantly greater degree of cognitive impairment, as judged on mean MMS scores. On the basis of this finding, O'Keane and Dinan suggested that orofacial dyskinesia might prove to be a useful indicator of the severity of intellectual decline in patients with Alzheimer's disease.

Acute Extrapyramidal Side Effects: Parkinsonism and Akathisia

Susceptibility to acute drug-induced extrapyramidal side effects as a predictor of tardive dyskinesia was first suggested by Crane in 1972⁹⁰. He considered that tardive dyskinesia was more likely to emerge in patients who had developed parkinsonism as an acute side effect of antipsychotic drug treatment than in those who had not.

Based on clinical observation, Chouinard *et al.*⁹¹ suggested that patients with tremor or akathisia, which they described as "hyperkinetic" symptoms of parkinsonism, were more likely to manifest tardive dyskinesia than patients with "hypokinetic" symptoms, such as bradykinesia and rigidity. De Veugh-Geiss *et al.*⁹² elaborated on this idea, suggesting that in some cases akathisia represented a stage in a progression from parkinsonism to the development of orofacial and trunk and limb dyskinesia. Consistent with this notion, Barnes and Braude⁹³ reported two

cases where the appearance of akathisia at the beginning of drug therapy, which then persisted despite the reduction of drug dosage to maintenance levels, seemed to herald the early onset of tardive dyskinesia.

Thus, a variety of study findings and case reports suggest that patients presenting with symptoms of parkinsonism and acute akathisia are more likely to manifest tardive dyskinesia. More convincing evidence is available from a prospective study of risk factors for tardive dyskinesia carried out by Kane *et al.*¹¹. Analysis of the data from the first 5 years of their follow-up study suggested that a history of early, clinically significant parkinsonism indicated susceptibility to the subsequent development of tardive dyskinesia, particularly for those patients developing tardive dyskinesia within the first 2 years of exposure to antipsychotic drugs. The patients in this study were relatively young, the mean age of the 800 patients originally recruited being 27 years. Further supportive evidence was provided by the prospective studies of elderly patients conducted by these investigators^{26,27}. Individuals exhibiting signs of acute extrapyramidal side effects, i.e. parkinsonism and akathisia, early in treatment with antipsychotic drugs showed a greater vulnerability to tardive dyskinesia.

If susceptibility to akathisia is associated with susceptibility to tardive dyskinesia, it might be expected that the two conditions would commonly coexist in patients chronically treated with antipsychotics. Several studies have found this to be the case^{13,94}. For example, Barnes and Braude⁹⁵ reported that 39% of schizophrenic outpatients with chronic akathisia also had orofacial dyskinesia. Further, out of 52 cases of chronic akathisia, Burke *et al.*⁹⁶ found that all but one had either orofacial dyskinesia (63%), tardive dystonia (8%) or both (27%). Dufresne and Wagner⁹⁷ also reported an association between the two conditions. In 33 chronic schizophrenic patients, these investigators found that the mean AIMS score (see earlier) was significantly higher in those patients with akathisia compared with those without the condition.

SPONTANEOUS DYSKINESIA

A condition described as spontaneous or idiopathic orofacial dyskinesia has been noted in psychiatric inpatients, predominantly elderly, who have never received antipsychotic drugs^{8,98-100}. The prevalence figures in the literature vary from 0% to around 37%, but it is not always clear whether the hyperkinetic movements disorders reported represent senile chorea, spontaneous orofacial dyskinesia or senile dyskinesia^{101,102}. These spontaneous orofacial movements show an increase in incidence with age. In a sample of 661 patients, Klawans and Barr¹⁰³ found a prevalence of 0.8% between the ages of 50 and 59, 6% between 60 and 69 years, and nearly 8% in patients between 70 to 79 years.

Crane¹⁰⁴ has stated that "chronicity of disease and/or institutionalization with attendant emotional and physical deprivation" may be responsible for these abnormal movements, being a common factor in both drug-treated and non-drug-treated psychiatric inpatients. Nevertheless, Altrocchi¹⁰⁵ reported two outpatients presenting with spontaneous orofacial dyskinesia, and this condition has also been observed as a relatively rare phenomenon in non-psychiatric, drug-free residents in old-age homes^{48,49}. It has been estimated that this condition is present in some 2% of the geriatric home population^{106,107}, although Varga *et al.*¹⁰⁸ surveyed a population of elderly people never exposed to antipsychotic drugs and discovered that 10% had clear evidence of oral dyskinesia. In a similar sample, Bourgeois *et al.*^{109,110} found that 18% exhibited "buccolinguofacial dyskinesia". These cases of spontaneous orofacial dyskinesia are generally considered to be indistinguishable from orofacial tardive dyskinesia, although

subtle differences in the distribution of movements have been detected between elderly psychiatric patients who were receiving antipsychotic drugs and normal elderly individuals who were drug-naïve¹¹¹.

Blowers *et al.*¹¹² surveyed 500 elderly residents in local authority homes using the AIMS. Out of the 378 individuals who had never received antipsychotic drugs, 50 (15%) were diagnosed as having "tardive dyskinesia" as they scored 3 or more on the AIMS global assessment scale. However, the AIMS may have limitations as a diagnostic scale^{113,114}.

These data generally suggest that elderly individuals are at a greater risk of spontaneous dyskinesias and abnormal involuntary movements related to various neurological and medical conditions independent of antipsychotic drugs. Nevertheless, surveys of normal elderly subjects suggest that spontaneous dyskinesias are relatively uncommon in the absence of brain disease or dysfunction.

SUMMARY

Elderly individuals would seem to be particularly vulnerable to the development of tardive dyskinesia when administered antipsychotic drugs, with females and those patients with dementia, other neurodegenerative conditions and organic brain syndromes being perhaps especially at risk. A diagnosis of affective disorder and the development of acute extrapyramidal side effects, such as parkinsonism and akathisia, when starting antipsychotic medication should also serve as warnings of a higher risk for tardive dyskinesia observed in elderly populations. Preliminary data suggest that use of the newer atypical antipsychotics, rather than conventional antipsychotics, in the elderly may substantially reduce the proportion developing tardive dyskinesia. A proportion of the orofacial dyskinesia observed in elderly populations, between 5% and 15% according to the majority of studies, will be spontaneous dyskinesia of the elderly. This is a condition virtually indistinguishable from tardive dyskinesia.

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Part H

Neuroses

Nosology and Classification of Neurotic Disorders

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The grouping of a variety of psychopathologic entities under the heading of “neurotic disorders” represents the current attempt to solve a nosological tangle dating back further than the time of Sigmund Freud. The contemporary categorization is controversial and less than satisfying to many. A brief look at its historical roots offers some explanation of the sense behind the current ICD-10 and DSM-IV structures.

HISTORY OF THE CLASSIFICATION OF THE NEUROSES

It was Hippolyte-Marie Bernheim who introduced the term “psychoneurosis” for hysteria and allied conditions. Freud, who studied with Bernheim, differentiated the “actual neuroses” (including neurasthenia, anxiety neurosis and hypochondria) from the psychoneuroses. The latter category included not only the “transference neuroses”, such as hysteria and obsessive neurosis, but also the psychoses (paraphrenia, schizophrenia, paranoia and manic depression), perversions and neurotic character. Both types of neurosis were related to sexual disturbance; the actual neuroses were direct somatic consequences of a noxious physical influence resulting from misdirected sexual energy; the psychoneuroses were caused by unconscious conflict between instinctual and counter-instinctual forces¹. Although Freud’s thinking on the precise nature of the mental etiology of the psychoneuroses changed over the years from about 1894 to 1906, he remained consistent in his stance that the psychoneuroses were defined by their etiology rather than by their phenomenology². Eventually, the psychoses were classified independently, consistent with the views of Kraepelin.

In 1952, the American Psychiatric Association published the first *Diagnostic and Statistical Manual of Mental Disorders* (DSM-I). It identified the subtypes of psychoneurotic disorders as anxiety, dissociative, conversion, phobic, obsessive-compulsive, and depressive reactions³. By the mid-1960s, the major diagnostic schemata, ICD-8 and DSM-II, codified the selection of the descriptive framework for identifying the neuroses. Anxiety was seen as the chief characteristic, whether felt and expressed directly or diverted unconsciously into other symptoms. The neuroses were also grouped by severity; they were more specifically symptomatic than the personality disorders, but entailed no gross distortion or impairment of reality testing, as in the psychoses. Categories included in the 300-code section were anxiety, hysterical, phobic, obsessive-compulsive, depressive, neurasthenic, depersonalization and hypochondriacal neuroses.

Transient situational disturbances constituted their own category (code 307)⁴.

The descriptive focus was emphasized in ICD-9 with the substitution of the term “neurotic disorders” for “neuroses”, although the categorization was not significantly modified⁵. In 1980, the American Psychiatric Association published the DSM-III⁶, which took a substantial leap towards atheoretical descriptive diagnosis by adopting empirically validated criteria based on research diagnostic criteria. One of the most controversial changes was the elimination of the entire class of neuroses. The neurotic disorders were included in the affective, anxiety, somatoform, dissociative and psychosexual disorders⁷. The grouping by severity was abandoned in favor of clusters based on similarity of features. The diagnostic entities retained numerical codes compatible with ICD-9 and ICD-9-CM.

The other revolutionary change introduced with DSM-III was the use of a multiaxial system for diagnosis. Under this scheme, personality disorders, which often predispose individuals to the development of specific neurotic (and other) syndromes, were relegated to a separate and parallel Axis II. The 1987 revision, DSM-III-R, changed some names and criteria but retained the same hierarchy of the neurotic disorders and the same multiaxial formula⁸. DSM-IV, published in 1994, was a more substantive revision overall than DSM-III-R, with only a few changes relevant to the neurotic disorders. The diagnosis of Acute Stress Disorder was added for compatibility with ICD-10. Dissociative Identity Disorder was added to Axis I to replace Multiple Personality Disorder, which was removed from Axis II. Simple Phobia was renamed Specific Phobia for compatibility with ICD-10⁹.

Neurotic Disorders in ICD-10 and DSM-IV

The creators of ICD-10 were faced with a formidable challenge, as this version, unlike its nine predecessors, was to be designed as the last of the series to be scheduled for regular revisions¹⁰. It therefore had to contain a format that would allow for flexibility in minor revisions while establishing a more permanent structure than versions 1–9. While following the lead of the phenomenological school in separating out a major category for mood disorders that includes both psychotic and neurotic levels of severity, it retains the major classification of neurotic disorders, including somatoform disorders and stress-related disorders¹¹. This grouping solves the objection that the term “neurosis” groups together entities which could be better classified, e.g. by placing neurotic severities of depression together with other mood

Table 97.1 Relative positions of neurotic disorders in the diagnostic schemata

ICD-10	DSM-IV
F0 Organic mental disorders	Disorders usually first evident in infancy, childhood or adolescence
F1 Mental and behavioral disorders due to psychoactive substance uses	Delirium; dementia and amnesic and other cognitive disorders
F2 Schizophrenia, schizotypal states and delusional disorders	Mental disorders due to a general medical condition not elsewhere classified
F3 Mood disorders	Substance-related disorders
F4 NEUROTIC, STRESS-RELATED AND SOMATOFORM DISORDERS	Schizophrenia and other psychotic disorders
F5 Behavioral syndromes and mental disorders associated with physiological dysfunction and hormonal changes	Mood disorders
F6 Abnormalities of adult personality and behavior	ANXIETY DISORDERS
F7 Mental retardation	SOMATOFORM DISORDERS
F8 Developmental disorders	DISSOCIATIVE DISORDERS
F9 Behavioral and emotional disorders with onset usually occurring in childhood or adolescence	Factitious disorders
	Sexual and gender identity disorders
	Eating disorders
	Sleep disorders
	Impulse-control disorders not elsewhere classified

disorders, rather than with anxiety disorders. It does, however, retain the historical commonality that traces to Freud's original stress on the etiologic similarity of the psychoneuroses, acknowledging the current state of scientific knowledge that is, at best, ambiguous concerning the etiology of these disorders^{12,13}.

Table 97.1 compares the relative positions of the neurotic disorders in ICD-10 and Axis I of DSM-IV. The alphanumeric organization of the International Classification requires the constraint of all mental disorders to 10 major categories. DSM-IV, under no such limitation, separates out anxiety, somatoform and dissociative disorders but keeps them within the same gradient of severity between mood disorders and sexual disorders. Adjustment disorders are removed to a position implying less severity, as well as an implied direction that higher-ranking diagnoses are to be made or eliminated first. Personality disorders, of course, are assigned to Axis II.

Under the category of the neurotic disorders, the international and American systems differ in their organization, as outlined in Table 97.2. DSM-IV groups the phobic disorders, obsessive-compulsive disorder, post-traumatic stress disorder and generalized anxiety disorder together as anxiety disorders. ICD-10 separates phobic disorders, anxiety disorders, and obsessive-compulsive disorders. Post-traumatic stress disorders are classified with adjustment disorders. Both schemes separate dissociative and somatoform disorders. ICD-10 groups conversion disorder with dissociative states, consistent with the historical, etiologically-based classification of the hysterias. DSM-IV combines it with the somatoform disorders, based on their phenomenological similarities. ICD-10 retains the diagnosis of neurasthenia; while DSM-III-R refers the clinician to dysthymia, categorized unequivocally as a mood disorder, DSM-IV eliminates the term entirely^{8,9,11}.

DIAGNOSTIC FEATURES OF NEUROTIC DISORDERS

While each of the major categories of neurotic disorders is described in the chapters that follow, the clinical features of the seven ICD-10 groupings are presented here for overview and comparison¹¹.

Phobic disorders are a set of disorders in which anxiety is invoked only, or predominantly, by well-defined situations that are not in themselves dangerous. By definition, the object of the fear is external to the individual, so that fears of bodily processes are more appropriately relegated to the category of somatoform

disorders. The feared objects are characteristically avoided, and anticipatory anxiety is common.

Other anxiety disorders are those in which anxiety is the major symptom but which are not restricted to specific situations. They include panic disorder, generalized anxiety disorder, and mixed anxiety and depressive disorder. The latter is reserved for cases where symptoms of both are present, neither is predominant, and the depression is not severe enough to be classified under mild depressive disorder.

Obsessive-compulsive disorder is characterized by recurrent obsessional thoughts or compulsive acts, or both. These thoughts and acts are subjectively distressing. Subjective anxiety is usually present and depressive features are common.

Reactions to severe stress and adjustment disorders represent a unique category, in that the component disorders are identified on the grounds of both symptomatology and causation. In these disorders, anxiety follows an exceptionally stressful life event or a significant life change. While psychosocial stressors may precipitate a wide variety of psychiatric syndromes, they are elsewhere neither necessary nor sufficient to explain the occurrence and form of the disorder. The stress and adjustment disorders, however, are seen as arising as a direct consequence of the trauma or life change.

Dissociative disorder is a group of entities that share a partial or complete loss of the normal integration between memory of the past, awareness of identity and immediate sensations, and control of bodily movements. It is presumed that, in these disorders, the ability to exert conscious and selective control over memory, sensation or bodily function is impaired.

Somatoform disorders are those in which physical symptoms are repeatedly presented with requests for investigation or treatment, in spite of the absence of physical findings to substantiate the perception. Compared with patients who suffer from psychogenic movement or sensory disorders, those with somatoform disorders will demand attention and usually resent physicians who fail to believe in the physical nature of their illnesses. Even when the onset of symptoms is temporally related to a stressful life event, or when external manifestations of depression or anxiety are obvious to others, these patients will frequently resist speculation about psychological causation.

Other neurotic disorders feature two clinical entities, neurasthenia and depersonalization-derealization syndrome. The former, recalling the pre-Freudian nomenclature, is a controversial category in contemporary psychiatry. Its main feature is fatigue, which may occur upon either mental or physical exercise. The diagnosis is to be made only after depressive and anxiety

Table 97.2 Classification of neurotic disorders

ICD-10	DSM-IV
F40 <i>Phobic disorder</i>	<i>Anxiety disorders</i>
40.0 Agoraphobia	300.21 Panic disorder with agoraphobia
.00 Without panic disorder	300.01 Panic disorder without agoraphobia
.01 With panic disorder	300.22 Agoraphobia without history of panic disorder
40.1 Social phobias	300.29 Specific phobia
40.2 Specific (isolated) phobias	300.23 Social phobia
F41 <i>Other anxiety disorders</i>	300.3 Obsessive-compulsive disorder
41.0 Panic disorder	309.89 Post-traumatic stress disorder
41.1 Generalized anxiety disorder	308.3 Acute stress disorder
41.2 Mixed anxiety and depressive disorder	300.02 Generalized anxiety disorder
F42 <i>Obsessive-compulsive disorder</i>	<i>Somatoform disorders</i>
42.0 Predominantly obsessional thoughts	300.81 Somatization disorder
42.1 Predominantly compulsive acts	300.11 Conversion disorder
F43 <i>Reaction to severe stress and adjustment disorders</i>	307 Pain disorder
43.0 Acute stress reaction	300.7 Hypochondriasis
43.1 Post-traumatic stress disorder	300.7 Body dysmorphic disorder
43.2 Adjustment disorder	<i>Dissociative disorders</i>
.20 Brief depressive reaction	300.12 Dissociative amnesia
.21 Prolonged depressive reaction	300.13 Dissociative fugue
.22 With predominant disturbance of other emotions	300.14 Dissociative identity disorder
.23 With predominant disturbance of conduct	300.6 Depersonalization disorder
.24 With mixed disturbance of emotions and conduct	<i>Adjustment disorder</i>
F44 <i>Dissociative and conversion disorder</i>	309.0 With depressed mood
44.0 Psychogenic amnesia	309.24 With anxiety
44.1 Psychogenic fugue	309.40 With mixed disturbances of emotions and conduct
44.2 Psychogenic stupor	309.28 With mixed anxiety and depressed mood
44.3 Trance and possession states	309.3 With disturbance of conduct
44.4 Psychogenic movement disorders	
44.5 Psychogenic convulsions	
44.6 Psychogenic anaesthesia and sensory loss	
F45 <i>Somatoform disorders</i>	
45.0 Multiple somatization disorder	
45.1 Undifferentiated somatoform disorder	
45.2 Hypochondriacal syndrome	
45.3 Psychogenic autonomic dysfunction	
45.4 Psychogenic pain	
F48 <i>Other neurotic disorders</i>	
48.0 Neurasthenia	
48.1 Depersonalization–derealization syndrome	

Special Considerations in Geriatric Patients

Anxiety, both as a symptom and as a disorder, is common among the elderly, but not remarkably more or less so than at other ages. The nature of worry and its clinical manifestations, however, change with increasing age. The intricate relationships among psychosocial stress, physical illness, depression and anxiety in late life make the recognition, diagnosis and classification of neurotic disorders in the elderly quite complex^{14–16}.

The clinician can usually compare the fears and concerns of a younger patient against those of his/her own peers and arrive at a credible assessment of whether or not the anxieties are pathological. The aged, however have different fears; they worry about physical illness, crime, institutionalization, financial disaster, senility and physical dependency. It is often hard for the younger physician to determine whether the subjective interpretation of events, or the anticipation of future events, is in the realm of clinical anxiety or constitutes adaptive concern. Anxiety results from feelings of vulnerability, and the elderly are truly vulnerable in many ways. It is no easy task to diagnose agoraphobia in an 80-year-old person whose fear of crime in her neighborhood may exceed its statistical likelihood. The clinician walks a fine line between pathologizing a healthy response and failing to recognize neurotic dysfunction¹⁴.

Physical illnesses with psychiatric manifestations increase in prevalence with age, as does the need to take medications with emotional or behavioral side effects. Emphysema, for example, may produce features indistinguishable from those of panic disorder. Hyperthyroidism is commonly accompanied by symptoms resembling those of generalized anxiety disorder. Further, the guiding symptom profiles for the underlying disorders may be absent or muted in the aging person. “Silent myocardial infarction” and afebrile pneumonia are fairly common. Finally, the elderly consume significantly more medication than do younger people and exhibit psychiatric side effects at lower doses and serum levels. Bronchodilators may produce the symptoms of many anxiety states; recommended doses of over-the-counter medications for sleep or colds may induce presentations resembling dissociative states¹⁶.

As could be expected, the diagnosis of somatoform disorders and hypochondriasis is particularly complicated in the elderly. Somatic complaints are common. To some extent, the somatic presentation of emotional disorders is a sociocultural cohort phenomenon. The generation of people over 70 in the 2000s, for example, grew up in the 1940s and earlier. At that time, words such as “depression” and “anxiety” were not commonplace parts of everyday conversation. Emotional introspection was not culturally normative. Thus, the older person who complains today of having “butterflies in my stomach” may be aware of the physical concomitants of anxiety, but not of the emotional state underlying it. The clinician must “translate” somatically-phrased complaints to help determine the affective condition.

Furthermore, the increase in prevalence of almost all physical illnesses with age confounds the determination of pathological perception and behavior, necessary for making diagnoses of somatoform disorders. Both DSM-IV and ICD-10 leave room for a subjective judgment of whether the presence of physical symptoms is sufficient to explain the intensity of the patient’s response. There are no objective grounds for deciding when a complaint of abdominal pain constitutes a somatoform disorder in a person with concurrent emphysema, arthritis and congestive heart failure^{14,17}.

While the delineation of the diagnosis and treatment of post-traumatic stress disorder (PTSD) followed the societal impact of returning Vietnam War veterans, the syndrome is not uncommon in older individuals. The trauma may have been a different war (World War II or Korea), a natural disaster, or a personal event

disorders have been ruled out. Depersonalization–derealization syndrome is a rare disorder in which the patient feels that his/her own mental activity, body or surroundings are changed in quality so as to be unreal or remote. This phenomenon is more commonly observed as a feature of depression, phobias, obsessive-compulsive disorder and some psychoses.

such as physical assault. Symptoms may occur early and continue for decades. In many cases, symptoms may not even be manifest until many years after the traumatic event. Often, the symptoms of delayed PTSD are precipitated by a psychologically reminiscent contemporary event; a concentration camp survivor may not experience stress-related symptoms overtly until becoming widowed and being institutionalized half a century later^{14,15}.

SUMMARY

The current classification of neurotic disorders is the most recent step in the evolution of the nosological understanding of a diverse group of syndromes. The ICD-10 grouping represents a compromise between the phenomenological grouping of neurotic conditions on a scale of severity between healthy and psychotic function, and the etiological clustering of disorders presumed to arise from internal conflicts and vulnerabilities to external stressors. The ambiguities inherent in this system reflect the incomplete state of knowledge about the etiologies of the constituent conditions. The North American schema of DSM-IV sets aside questions of etiology, except in the case of adjustment disorders, and relies on ostensibly atheoretical phenomenological criteria.

Although the diagnostic criteria are technically independent of the age of the patient in both systems, aging affects the presentation of many of these disorders and makes clinical diagnosis challenging. The multiple biological and social stressors of late life blur the distinction between “normal” and “pathological” responses to these threats. Physical illnesses, which increase in frequency with aging, may produce clinical symptoms easily mistaken for neurotic anxiety. The prevalence of somatic pathology forces subjective judgments about the presence of somatoform and conversion disorders. Chronic and delayed stress reactions are clinically distinct from the acute forms seen in younger individuals.

The lack of clarity in the classification of these disorders, however, is probably less a manifestation of the shortcomings of the nosological systems than a reflection of the complicated function of the human mind. The neurotic disorders, as well as current science can determine, are a product not of brain disease but of human response to a complicated and stressful world. Simplicity in their nosology would belie the challenges they pose to patients and clinicians.

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Epidemiology of Neurotic Disorders

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Neurotic disorders are grouped as a special category (the 300 group) in ICD-9¹. They are mental disorders without any demonstrable organic basis, in which the patient may have considerable insight, and has unimpaired reality testing, i.e. he/she usually does not confuse a morbid subjective experience and fantasies with external reality. Included within the neurotic disorders are anxiety states (both generalized anxiety and panic disorder), hysteria (including conversion reactions), phobic states, obsessive-compulsive disorder, neurotic depression, neurasthenia and depersonalization syndrome.

*The Diagnostic and Statistical Manual of Mental Disorders*² of the American Psychiatric Association, in contrast, does not recognize the neurotic disorders as a distinct entity. Rather, all mood disorders (including the more severe depressive disorders, such as bipolar disorder) are classified in the same category with dysthymia or neurotic depression. Anxiety disorders in DSM-IV include generalized anxiety, panic disorder, phobic disorders and obsessive-compulsive disorders. There is no category within DSM-IV for neurasthenia.

As the epidemiology of the mood disorders will be covered elsewhere, in this chapter current knowledge regarding the epidemiology of the neurotic disorders will be reviewed with especial attention to anxiety. All extant studies that provide useful estimates by age are prevalence studies. Few investigators have concentrated specifically on risk factors by age. In addition, a Medline search of prevalence and incidence studies of generalized anxiety, panic, phobic and obsessive-compulsive disorders in late life since 1996 uncovered only one study. Therefore, the focus of this chapter will be predominantly upon estimates of prevalence of the neurotic disorders by age. Much of the review will feature the Epidemiologic Catchment Area (ECA) studies in the USA.

HISTORICAL REVIEW

At least two early studies provide estimates of the prevalence of neurotic disorders by age. Leighton *et al.*³, classifying “psycho-neurotic” disorders according to DSM-I in the Sterling County study, found that women experienced a peak in psychoneurotic disorders in the 40–49 decade, while the largest percentage of male psychoneurotics were found in the 60–69 decade. For both groups, there was a significant decrease in prevalence in subjects over the age of 70. Pasamanic *et al.*⁴ reported an overall prevalence of psychoneurotic disorders of 5.3%. The rates were lowest for subjects under the age of 15 but remained constant through most of adult life. For the 65+ age group, the prevalence estimate was 7%. These investigators of the epidemiology of psychoneurotic disorders found little evidence

of an increased prevalence by age but neither did they find evidence for a decrease. In these studies, however, the number of subjects aged 65+ was generally too small to make accurate estimates.

Kay *et al.*⁵, in their survey of older persons in Newcastle upon Tyne, which included both institutional cases and those individuals living in the community, found a prevalence of 10.2% for moderate to severe forms of neuroses in older persons. They did not compare the prevalence of neuroses by age. Bergmann⁶ found that 11% of a community sample of 300 persons reported developing a neurosis after the age of 60. He also noted the association between neurotic disorders and physical health problems. Each of the above studies aggregated all psychoneurotic disorders and did not exclude depression.

Regarding specific neurotic disorders, few investigators have reported prevalence estimates by age (except for the Epidemiologic Catchment Area studies). The exception is generalized anxiety, one of the more common disorders regardless of age. Murphy⁷ and colleagues estimated the prevalence of both anxiety disorders and conversion disorders in individuals less than 45 years of age vs. those 45 years of age and older in the Sterling County study described above. For these estimates, however, they applied diagnostic criteria similar to those used in the third edition of the *Diagnostic and Statistical Manual*² rather than the more inclusive category of psychoneurosis. The prevalence of generalized anxiety was 2.9% overall. In males the prevalence was 2.1% under the age of 45 but less than 1% in individuals 45 years of age and older. The prevalence of generalized anxiety in females 45 years of age and older was 2% compared to 6% for those under the age of 45.

Warheit *et al.*⁸, in a study of community-dwelling adults in northern Florida, used a self-rating anxiety scale and found that 14.6% of the sample had significant symptoms of anxiety. Blacks, females, the elderly, those in the lower socioeconomic levels and those separated, widowed and divorced actually had the highest scores (in contrast to the ECA study). In a previous study using the Schedule of Affective Disorders and Schizophrenia–Lifetime Version (SADS-L), the overall prevalence of generalized anxiety was 2.5% and was slightly more common in middle-aged and younger women than in older adults.

Copeland *et al.*⁹ in a survey of 1070 elderly persons (65+) living in Liverpool, using the Geriatric Mental State (GMS), found the overall levels of neurotic disorders in the elderly to be 2.4%. Anxiety disorder was the most prevalent of the neurotic disorders, with phobic disorder being the second most prevalent (see below). Hypochondriasis and obsessive-compulsive disorders were found in approximately 0.5% and 0.2%, respectively. During a 3 year follow-up¹⁰, the incidence was 0.44%/year. Women were more

Table 98.1 One year prevalence rates (%) of generalized anxiety (different exclusionary criteria) by age. ECA study¹²

Age	Generalized anxiety/no exclusions	Generalized anxiety/no panic or major depression	Generalized anxiety/no DSM diagnosis
< 30	4.83	3.51	2.35
30–44	3.58	2.12	1.46
45–64	3.74	2.81	1.75
65+	2.22	1.92	1.05

Table 98.2 Cumulative prevalence by age of onset of cases of generalized anxiety (no panic or major depression). ECA study¹²

By age	Cumulative prevalence (%)
19	20.6
24	40.1
29	56.5
44	79.5
64	97.0

Table 98.3 Use of inpatient and outpatient general health services by age and diagnosis of generalized anxiety. ECA study¹²

Age	Outpatient use		Inpatient use	
	Generalized anxiety (%)	No generalized anxiety (%)	Generalized anxiety (%)	No generalized anxiety (%)
45–64	37.0	52.8	10.3	5.7
65+	53.5	60.8	38.4	12.0

likely to become cases than men. Many cases did remit during the 3 years between interviews, but most original cases continued to experience some symptoms at follow-up.

FINDINGS FROM THE EPIDEMIOLOGICAL CATCHMENT AREA SURVEY

The most detailed estimates of specific neurotic disorders are derived from the ECA surveys in the USA. The National Institute of Mental Health Multi-Site Epidemiologic Catchment Area (ECA) Program was a collaborative study that combined community and institutional surveys of five communities in the USA—New Haven, CT; Baltimore, MD; Durham, NC; St Louis, MO; and Los Angeles, CA. The large overall sample (over 18 000 subjects), coupled with oversamples of older persons in three of the five sites, provided the most comprehensive estimates of the

prevalence of specific anxiety disorders among community-dwelling elders from any extant study. The instrument used to establish cases was the Diagnostic Interview Schedule (DIS).

The prevalence of generalized anxiety disorder at three of the ECA sites by age is presented in Table 98.1. The patterns presented hold for both males and females and for Whites, Blacks and Hispanics (although the patterns for male Hispanics are less obvious than for the other age, sex and race groups). In all groups, the prevalence for generalized anxiety is relatively high, but is lower in the 65+ age group than for other ages. Prevalence is presented when the symptoms of generalized anxiety are present, regardless of the symptoms of other disorders, when generalized anxiety is present without evidence of panic or major depression, and when generalized anxiety is diagnosed with no other DIS/DSM-III disorders. The patterns by age are the same regardless. In a study from Liverpool, Copeland *et al.*⁹ found the prevalence of cases of anxiety among females 65+ years of age to be 1.52%, yet nearly 16% of males and females were classified as subcases. Generalized anxiety is more frequent among persons with dementing disorders than without.

In a further analysis of the ECA data, the age of onset of generalized anxiety is presented in Table 98.2 for Durham, NC. Virtually all cases of generalized anxiety have their onset before the age of 65 in this community sample. Age of onset is evenly distributed across the life cycle except for individuals aged 65+.

In Table 98.3, the use of inpatient and outpatient general health service, with and without generalized anxiety disorder, are compared by age (persons 65+ years of age and persons 45–64). Older persons are more likely to report use of inpatient physical health services if they report a current episode of generalized anxiety. In contrast, older persons who suffer generalized anxiety are no more likely to use outpatient services. The trend in older persons is similar to trends in younger persons for both inpatient and outpatient use.

In Table 98.4, data are presented on other selected neurotic disorders by age and sex from the ECA study. In most cases, for both sexes, the prevalence of neurotic disorders decreases with age. Age differences in the rates of phobic disorder are not as pronounced as those seen for other disorders. Older persons have the lowest rates of panic disorder of any age group, whereas persons in the 30–44 age group have the highest rates. This trend occurs for Whites and Blacks but not for Hispanics. Not only do older persons appear to experience a lower prevalence of panic disorder in late life currently, they also appear to have a lower lifetime prevalence of panic disorder. This lower lifetime prevalence could be explained by the fact that persons with panic disorders are less likely to reach old age. In addition, the cohort phenomenon, which is described frequently throughout this book (i.e. persons in late life currently have been uniquely protected against a number of psychiatric disorders) may be operative for panic disorders as well. One must also consider, however, the possibility that older persons in these community surveys fail to recall episodes of panic in the distant past because they have not experienced them recently or they may find such episodes embarrassing to report.

Table 98.4 One year prevalence rates (%) of selected neurotic disorders by age and gender

Disorder	18–29		30–44		45–64		65+	
	Male	Female	Male	Female	Male	Female	Male	Female
Phobic disorder ¹³	6.5	13.4	6.1	16.1	6.7	11.6	4.9	8.8
Panic disorder ¹³	0.6	1.1	6.7	1.9	0.7	1.1	0.04	0.4
Obsessive-compulsive disorder ¹⁴	1.8	2.6	1.9	2.2	0.8	1.2	0.8	0.9
Hypochondriasis ⁹							0.5	0.5

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Stress, Coping and Social Support

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While a multitude of studies have examined the effects of social and environmental stressors, coping behaviors and social support on psychological distress, relatively little is known about their effects on neurotic disorders in the elderly. The effects of stress, support and coping on depression have been studied extensively in the general population and among those aged 65. A smaller number of studies examines their effects on the anxiety disorders in the general population. Very little work has been done with neurotic disorders as the outcome of interest among older persons. Recent reviewers conclude that the anxiety and panic disorders among the elderly have received little attention and that a systematic examination of the risk factors associated with late-life anxiety disorders has barely begun¹⁻³. This is the case despite the fact that they are the most prevalent psychiatric conditions among the elderly, as they are among younger persons. Methodological problems involved with defining and operationalizing the anxiety disorders, disentangling anxiety from depression, and the transience of some symptom states account, in part, for the lack of attention they have received in epidemiologic studies^{3,4}.

Since the overwhelming majority of relevant studies deal with depression rather than the neurotic disorders, we will use these to examine the rationale and evidence for epidemiologic models linking stress, coping and support to psychiatric symptoms and disorder. Next, we will review a smaller but growing number of studies that have begun to examine whether stress, support and coping affect the anxiety disorders in a manner similar to their effects on depression. Since all but three of these anxiety studies are based on general population surveys rather than samples of the elderly *per se*, we will address the degree to which these studies are consistent with a conclusion that similar effects of stress, coping and support on neurosis are present among the elderly. Finally, we will point out key unresolved issues and the practical implications of the studies reviewed.

STRESS, COPING, SOCIAL SUPPORT AND DEPRESSION

Stress

Stressors refer to life experiences that may be perceived as threatening and/or challenging. They include discrete "stressful life events", such as changes in finances, health or marital status. They also include more enduring or chronic problems with regard to income, health or other areas of life. Reviewers are in accord that there is consistent evidence that stress is

associated with an increased risk of psychiatric symptoms and disorder⁵⁻⁷. Findings for stress and depression are based on samples of older adults as well as the general population and include prospective studies where stress precedes the onset of symptoms⁸⁻¹⁰.

Stressful life events are most likely to have negative health consequences if they are perceived as unexpected and undesirable⁵. Negatively-evaluated changes in health, family and living situations, work and finances have been shown to be strongly and positively related to depressive symptoms and major depressive episodes in the general population and in samples of older individuals^{5,11,12}. Chronic stressors include poverty, deteriorated neighborhood conditions and ill health, which have been shown to predict both the onset of depression and the course of recovery¹³⁻¹⁷. While cross-sectional studies show an association between cognitive impairment and depression⁴⁻¹⁷, prospective studies report mixed and inconclusive results regarding whether dementia or cognitive impairment are risk factors for the onset or duration of depression¹⁷.

While older persons experience fewer potentially stressful life events¹⁸, they experience a "changing landscape of stressors"¹⁹, and are more likely to experience particular events that are strongly related to psychiatric morbidity²⁰. These include poor health and disability, widowhood, and the death of other friends and family members^{21,22}. While retirement *per se* is not associated with an increased risk of psychiatric disorder²³, a recent study²⁴ reports that driving cessation is strongly associated with an increased risk of depressive symptoms. Findings are mixed regarding whether stress associated with caring for a disabled person is a risk factor for depression. Initial studies of those (presumably more distressed) caregivers who sought services found high levels of depressive symptoms. A smaller number of community studies of caregiving and depression report inconsistent results¹⁷.

Coping

Coping refers to steps the individual takes to avoid, solve or minimize the impact of life problems²⁵. It serves two functions: problem solving and the regulation of emotions. Different coping strategies can have different consequences for psychological well-being generally, and for the impact of stress on well-being in particular. Rodin²⁶ contends that solving a problem without help from others may promote well-being by enhancing feelings of self-worth and personal control. Requesting help, on the other hand, may negatively affect well-being by generating interpersonal conflict if those asked are unwilling or unable to provide

assistance⁹. This is especially true for financial help. One's social network is typically made up of persons in similar economic circumstances, whose financial resources may already be limited. There is evidence that most older people value independence and prefer to resolve problems by themselves, rather than depend upon others²⁷. Systematic evidence for the impact of specific coping strategies on psychiatric morbidity is presently lacking. The number and variety of possible coping responses, together with the fact that assessments of appropriate coping behavior may vary across situations and social groups, has made research in this area difficult²⁰.

Social Support

Social support refers to a number of different aspects of social relations and includes: (a) social network: the size, stability, and structure of an individual's network of friends, relatives, and acquaintances; (b) social interaction: the presence and quantity of interaction with network members as well as organizational participation; (c) instrumental support: services and assistance provided by family and friends; and (d) perceived support: subjective satisfaction with one's social relationships and availability of support.

Two alternative models have guided most empirical studies of the relationship between social support and depression. With the "stress-buffering" model, a statistical interaction is hypothesized. The protective effect of support is expected to be at its maximum under conditions of stress and weaker when stress is absent^{28,29}. Given the presence of a potentially stressful event or experience, social support is thought to influence the degree to which the situation is appraised as threatening, and an individual's capacity to cope. The impact of stress on depression will therefore be strongest among those lacking adequate support. In the absence of stress, the availability of support is of less importance, and the relationship between support and depression is expected to be weaker. With the "main effect" model of stress, support and depression, high levels of social support are hypothesized to promote mental health at all levels of stress. From this perspective, the effects of stress and support are not interactive—the effect of each does not depend upon the level of the other^{30,31}.

Reviewers report that the protective effect of support on depression varies across its different dimensions in the general population and among older persons^{8,9,20,21}. Perceived support is most strongly and consistently protective for depression. There is also longitudinal evidence that the primary causal influence is from perceived support to depression, rather than the reverse. Findings for the protective effect of network size have been mostly negative. Amount of social interaction is associated with depression in several studies, but not with the onset of depression in longitudinal research. Received support can increase, as well as reduce, the risk of depression^{9,12}. Krause *et al.*⁹ reason that the receipt of assistance may reflect a failed attempt to solve a problem on one's own, and may be accompanied by hostility and resentment from those providing assistance. Reviewers are also in agreement that most (but not all) studies report a stress-by-support interaction consistent with the stress-buffering models^{11,20}. Positive findings for stress buffering are most often present for perceived support, and recent findings suggest that anticipated support—the belief that others stand ready and able to help if called upon—is especially critical, as it promotes effective coping and confidence that a problem can be solved^{9,13}.

According to Kahn and Antonucci's convoy metaphor for social support³², the size and composition of one's social network changes over the life course as individuals enter and leave a

variety of social roles (e.g. spouse, parent, employee). Social networks change composition later in life in response to changes in one's health and employment, and to impairment and death among one's age peers²⁰. For example, retirement can provide time to expand the scope of one's social participation, and even poor health, which limits some relationships, may enhance others as one's support network is mobilized to provide assistance³³. Findings are mixed regarding whether there is a net decrease in network size and frequency of contact in old age, allowing different reviewers to draw different conclusions. However, there is general agreement that aging is not a time of social isolation, and that most older people have a significant number of relationships^{20,33}. Studies of changes in social network and interaction after age 65 report considerable change, characterized by widely varying patterns of gains and losses rather than a trend toward isolation³³. While it is unclear how these specific changes affect the psychological health of older adults, the notion that old age is a time of psychologically debilitating isolation is clearly not supported.

STRESS, COPING, SOCIAL SUPPORT AND THE NEUROTIC DISORDERS

The proposition that stress, support and coping may affect neurosis is consistent with existing theories that suggest that symptoms of anxiety and panic disorders represent a dysfunctional response to potentially stressful environmental events¹⁰. While anxiety is adaptive in the face of potentially threatening or unpleasant events, the anxiety disorders are characterized by unjustifiably intense and morbid anxiety and panic². Endler's multidimensional interaction model of anxiety includes situational factors (stressors) and individual characteristics which interact to produce anxiety symptoms and disorder³⁴. Relevant individual characteristics include "trait anxiety"—a predisposition to react to stressors generally, or to particular stressors, with dysfunctionally high and persistent levels of anxiety. Individual traits also include differentially effective coping styles and behaviors. One's appraisal and use of available resources, such as social support, is incorporated as part of coping.

There is also reason to expect differences in how stress and support might relate to anxiety as opposed to depression. The anxiety disorders, which include agoraphobia (with and without panic attack), social and simple phobia, panic disorder, generalized anxiety disorder and obsessive-compulsive disorder (OCD), are considerably more complex and diverse than the subtypes of depression. This has led reviewers to call for research that considers these subcategories separately in examining the effects of stress, support, coping and other risk factors^{34,35}. The "multi-dimensional" aspect of Endler's model refers to the proposition that trait anxiety may be stressor-specific. Environmental danger might trigger anxiety only among those predisposed on this trait, while a symptomatic response to a job interview might be limited to those differently predisposed. The implication—that the effect of a stressor on anxiety would be greatly attenuated unless it is estimated separately for those with the corresponding trait anxiety—adds considerable complexity to the stress-support model. A related hypothesis—that stressors dealing with loss (of health, finances or social support) might result in sadness and depression, while stressors involving danger (severe future threat but not necessarily loss) might trigger anxiety—further complicates the picture³⁶.

Reviewers agree that most studies report a positive association between various stressors and one or another measure of anxiety^{34,35,37}. Investigators report both transient and long-term symptoms of anxiety and depression following exposure to

extreme experiences (e.g. combat missions and natural and man-made disasters)³⁴. Chronic stressors and stressful life events are also related to anxiety^{37,38} and negatively-evaluated stressful events have a stronger impact than the number of events *per se*⁴. Stressors included negative life events, physical decline, poor health, chronic financial stress, occupational stress and loss of a family member. Longitudinal findings show that various stressors precede the onset of anxiety^{4,36,39,40} and influence treatment outcome over time³⁸. There is also evidence that minor stressors (“daily hassles”) may play a particularly important role in the onset of generalized anxiety disorder^{41,42}. Three studies examined the stress-anxiety relationship in elderly populations and report various stressors, including stressful life events, health problems and loss of a loved one to be significant risk factors for phobic disorders, generalized anxiety disorder and the anxiety disorders generally^{3,4,43}.

Coping style and social support are related to one or another measure of anxiety in prior studies. Panic disorder has been found to be associated with less effective coping strategies in response to stress⁴⁴⁻⁴⁶. Those with panic disorder are more likely to use strategies involving escape, avoidance, wishful thinking and help-seeking, rather than focused problem-solving without help. Studies based on age-heterogeneous and elderly samples of the elderly report that external locus of control is positively related to anxiety^{3,47}. Compared to others, those with generalized anxiety disorder are reported to perceive the same stressors as more stressful and threatening³⁵. Several measures of social support have been found to be associated with anxiety symptoms and/or disorder. These include small network size, marital problems, not having a confidante, and loneliness^{3,38,39,48}. In one study, death of a network member was found to affect depression but not anxiety³⁹. Another study, which focused on stress-buffering, reports that a stress-by-support interaction effect is present for anxiety but weaker than the corresponding effect for depression⁴⁹.

CONCLUSION

While the above studies provide evidence that stress, coping and support affect the anxiety disorders as well as depression, they are limited in number and scope and do not permit any firm conclusions about the specific workings of the stress-vulnerability model derived from depression studies, or the more complex multidimensional model proposed by Endler³⁴. This is especially true with respect to the elderly population, where we were able to locate only three relevant studies. The extensive literature on depression suggests that the effects of stress, support and coping on psychiatric disorder do not vary with age. However, researchers have only begun to address this issue with respect to neurosis. We located two studies comparing the effects of stress on anxiety in different age groups. One reported that the effects of different stressors were mostly the same in older and younger populations, while a second reported age differences in the impact of stressful life events^{3,4}.

Perhaps the most critical gap in the literature reviewed here is the absence of systematic knowledge about differences in the effects of stress, coping and support (and risk factors generally) across the specific subcategories of neurosis. Important in its own right, this information is especially critical if consistent and inconsistent findings across studies using different anxiety measures are to contribute to a cumulative understanding. Only one study has examined the effects of the same measures of stress, coping and support on the specific subtypes of anxiety, and reports that partner loss and poor health more strongly related to panic disorder and OCD, while network size and the exchange of emotional support affect only the phobias³. Their cross-sectional

data and the specificity of the sample (older adults in Holland) make this study an initial step in what needs to be an ongoing process.

Findings are also sparse and inconsistent with respect to other proposed links between neurosis and the risk factors considered. The hypothesis that anxiety and depression may result from different stressors (threat vs. danger) received support in an initial test³⁶. Two subsequent studies report that most stressors affected both anxiety and depression, and that differences which did exist were not consistent with the threat vs. danger hypothesis^{38,39}. Endler's proposition³⁴, that particular stressors may trigger anxiety only among those with a congruent susceptibility, has yet to be examined in representative community samples. While there is evidence that social support is protective for the neurotic disorders, we do not know which dimensions of support are critical, whether support exerts a generalized protective effect or whether stress-buffering is operative. Finally, as discussed in Monroe and Wade³⁵, substantial co-morbidity between anxiety and depression make it imperative that we examine the role of depression in the relationships of stress, coping and support to anxiety.

While epidemiologic examination of the effects of stress, coping and support on the neurotic disorders is still in an initial stage, the research reviewed here is not without practical import. As Sheikh² has observed, potential side effects, drug interactions and non-compliance among the elderly make effective non-pharmaceutical therapies particularly attractive². In this regard, clinicians and researchers have reported that the success of exposure-based treatments for agoraphobia depends in part on the patient's marital relationship, and that including the spouse in therapy can be critical to successful treatment⁵⁰⁻⁵². Findings reviewed here—that marital satisfaction is protective for anxiety symptoms in addition to agoraphobia—suggest that attention to the marital relationship may improve psychological and behavioral treatments for other anxiety disorders as well. Findings that other forms of social support are also protective suggest that focusing on relationships in addition to the marital one may also be useful. Additional studies of onset and recovery, which systematically related marital and other social relationships to onset and recovery for specific disorders among the elderly, would be especially useful in this area.

The studies reviewed also have implications at the social structural level. There is strong and consistent evidence that inequalities in education, occupation and income are major determinants of the public's mental health, and that the least privileged members of society are at increased risk for health, financial and work-related stressors, which contribute to both anxiety and depression^{37,53}. The well-to-do, on the other hand, are less exposed and have more social and material reserves with which to overcome negative and unpredictable events. Treating mental disorder at the social as well as the individual level involves programs and policies designed to reduce inequality and/or reduce its impact on mental health. Among the elderly, improved coverage for mental health treatment under Medicare and Medicaid would enable the elderly, and particularly those with limited incomes, to obtain necessary and timely treatment²⁰. Recognizing health problems as a major risk factor for depression among the elderly, Jorm advocates “improved geriatric care” to reduce depression¹⁷. While findings for the impact of health on neurosis are not nearly so extensive as for depression, poor health is a consistent risk factor for one or another measure of anxiety. In this regard, evidence is accumulating that expanded Medicare coverage for prescription drugs, preventive care and other services typically covered by supplemental private insurance would substantially reduce health problems and disability among older persons, especially those with limited financial resources⁵⁴.

Evidence that low income persons enter old age in considerably worse health than others⁵⁵ indicates that improving the mental health of the elderly will also require attention to social inequality and its link to physical and mental health throughout the life course.

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Clinical Features of Anxiety Disorders

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The subjective sense of trepidation or dread about some future event that can motivate one person to stay at work late to complete an important project, can send another to the hospital with the belief that he/she is going into cardiac arrest. As a normal human emotion, anxiety has adaptive value in helping one prepare for, and possibly avoid, deleterious events. This emotion, however, can manifest pathologically if it becomes excessive, inappropriate or maladaptive. Such morbid or clinically significant anxiety can range from excessive worry about mundane concerns to experiencing intense episodes of fear (panic attacks) for no apparent reason. Clinically significant anxiety is usually manifested by a variety of cognitive, behavioral and physiological symptoms. Table 100.1 lists some examples of these multi-dimensional features. When assessing such symptoms, the clinician will query the patient with regards to the duration, intensity and course to determine whether the cluster of symptoms meet criteria for any of the specific anxiety disorders.

Over the last two decades, researchers have made great strides in furthering the understanding of the phenomenology, comorbidity and clinical course of anxiety disorders in the general population¹⁻⁴. However, research with the geriatric population is lagging, forcing clinicians to use knowledge gained from studies that more commonly study a younger age group. In addition to utilizing these empirical studies, those treating the anxious elderly must rely mostly on their own observations and anecdotal information, in addition to the "youth-biased" literature base⁵. Some have expressed concern that the diagnosis of anxiety states may be particularly difficult in the elderly because of the frequent co-morbidity of depression or medical illness⁶⁻⁷. In addition, concerns are being raised regarding the *Diagnostic and Statistical Manual of Mental Disorders*, 4th edn (DSM-IV) of the American Psychiatric Association⁸. The DSM criteria for the anxiety disorders may not always allow for the correct identification of the anxiety-disordered elderly because many older adults display a tendency to deny cognitive symptomatology and instead somatize their distress⁹. Despite these limitations, analyses of the Epidemiologic Catchment Area (ECA) data (using the DSM-III-R criteria) indicate that anxiety disorders can be diagnosed successfully. Finally, the risk of developing an anxiety disorder does not fade in late life. For example, in a large sample of the 65+ age group, individuals experienced a rather high 6 month prevalence of 19.7% for all anxiety disorders¹⁰.

This chapter presents the clinical features of various anxiety disorders based on the criteria set out in the DSM-IV. A list of the DSM-IV anxiety disorders appears in Table 100.2. Differential presentations and unique features of the anxiety disorders in the elderly will be discussed where appropriate.

PANIC DISORDER WITH/WITHOUT AGORAPHOBIA (PD/PDA)

Panic attacks are acute and discrete episodes of intense anxiety that result as a reaction to some perceived threat (emotional, environmental, etc.). The term "panic attack" is used when an individual experiences an intense and acute reaction to an internal or external cue, lasting between a few minutes and a half an hour. The physiological symptoms can include trembling, accelerated heart rate, sweating, shortness of breath, chest pain, dizziness, nausea and the sense that one is somehow detached from one's surroundings¹¹. For example, an individual might have been trapped in a crowd of people entering an underground subway system, and will describe feeling "sick to their stomach" when entering one. Another individual might even report high levels of acute anxiety at the mere sight of the stairs leading to the subway. A clinically significant degree of panic symptoms are documented after a review of the patient's history, revealing recurrent and unpredictable panic attacks that precede at least 1 month spent with anticipated worry over possible recurrence.

Diagnostically, one needs also to consider whether there is the presence of agoraphobia in relation to the panic attacks. Agoraphobia involves the persistent fear of being in a situation that results in a panic attack. Individuals suffering from agoraphobia will commonly stay inside their house all day long to ensure the avoidance of the feared situation. Some of the common examples of frightening situations include being caught in traffic on a bridge or freeway. When comparing young and older adults with panic disorder (PD), one of the factors that can affect the clinical presentation appears to be the age of onset. Phenomenologically, it appears that late-onset PD (LOPD, onset of PD at or after age 55) patients report fewer panic symptoms, less avoidance, and score lower on somatization measures than do early-onset PD (EOPD, onset of PD prior to age 55) patients.

AGORAPHOBIA WITHOUT HISTORY OF PANIC DISORDER (AWOPD)

The literature is scant regarding this relatively rare disorder. Its distinguishing feature from Panic Disorder with Agoraphobia is a fear of being in public places or situations from which escape might be difficult, even though there is the *absence* of a history of panic attacks. There is a possibility that, although these patients may not experience full-blown panic attacks, they might suffer from milder ones with only one or two symptoms (limited symptom attacks). It is thus possible that some of these patients

Table 100.1 Multidimensional symptoms of anxiety

Cognitive	Behavioral	Physiological
Nervousness	Hyperkinesis	Muscle tension
Apprehension	Repetitive motor acts	Chest tightness
Worry	Avoidance (e.g. certain places)	Palpitations
Fearfulness	Pressured speech	Hyperventilation
Irritability	Increased startle response	Paresthesias
Distractibility	Lightheadedness	Lightheadedness
	Sweating	Sweating
	Urinary frequency	Urinary frequency

are presenting with a variant of panic disorder. Multicenter studies suggest that AWOPD, GAD and Social Phobia are commonly co-morbid. Moreover, in general, AWOPD presents with worse global functioning than PD or PDA¹². This syndrome has not been studied in the elderly.

SOCIAL PHOBIA (SOCIAL ANXIETY DISORDER)

Social Anxiety Disorder (SAD) is defined by a persistent fear in one or more social situations marked by fears of performance, excessive scrutiny or of acting in a way that will be embarrassing or bring shame. Frequently, the fear is that of trembling, blushing or sweating profusely in social situations. Other common concerns are of saying something stupid or “babbling or talking funny”. Common examples include fears of public speaking, avoidance of dating, parties or other social gatherings. Social phobics typically experience marked anticipatory anxiety if they attempt to enter the phobic situation. SAD is associated with onset in early life—typically manifesting in adolescence. The two distinct subtypes, generalized and non-generalized, trigger different types of symptoms, course of illness, pathophysiology and response to treatment¹³. Although systematic studies of this disorder in the elderly are lacking, epidemiological data¹¹ indicate that it is chronic and persistent in old age. Common manifestations in old age include the inability to eat food in the presence of strangers and, especially in men, being unable to urinate in public lavatories. It is unlikely that an older adult will seek professional help with these complaints as primary. Although systematic studies of social phobia in older patients are lacking, our clinical experience suggests that eating or writing in public can be exceedingly difficult in older social phobics, exacerbated by the use of dentures or the presence of tremors. It is not uncommon to encounter social phobics who present with symptoms of panic disorder. Evidence suggests that this disorder is quite commonly co-morbid with panic disorder¹⁴.

SPECIFIC PHOBIA (FORMERLY SIMPLE PHOBIA)

The distinguishing feature of this disorder is a marked and persistent fear of a specific object or situation (other than a fear of experiencing a panic attack or a fear of social situations). Typically, the patient experiences immediate and intense distress on encountering the phobic stimulus, and recognizes that the fear is excessive and/or unreasonable. Further, the avoidance or anxious anticipation of encountering the phobic stimulus must interfere with the person's daily routine, occupational functioning or social life, or the individual is markedly distressed about having the phobia. The level of anxiety or fear usually varies as a function of both the degree of proximity to the phobic stimuli and the degree to which escape is limited. Examples of common phobias include fear of animals (dogs, snakes, insects, etc.), closed spaces (claustrophobia), flying or heights. There is

Table 100.2 DSM-IV anxiety disorders

● Panic disorder without agoraphobia (PD)
● Panic disorder with agoraphobia (PDA)
● Agoraphobia without history of panic disorder (AWOPD)
● Social phobia (social anxiety disorder, SAD)
● Specific phobia (formerly simple phobia)
● Obsessive-compulsive disorder (OCD)
● Acute stress disorder (ASD)
● Post-traumatic stress disorder (PTSD)
● Generalized anxiety disorder (GAD)
● Anxiety disorder not otherwise specified (ADNOS)
● Anxiety disorder due to a general medical condition
● Substance-induced anxiety disorder

frequent co-occurrence of Specific Phobia with PD and PDA. In the elderly, especially in urban settings, fear of crime seems to be particularly prevalent in the elderly population (although they are the least likely to be victimized). UK researcher Lindsay¹⁵ looked at elderly phobics and matched them for age and sex to case controls without history of phobic disorders, and found that in the elderly phobic disorders are associated with considerably higher psychiatric and medical morbidity. It also appears that, despite higher rates of contact among the phobic elderly with general practitioners compared to controls, only 1 in 60 of the phobic elderly in this study was receiving psychiatric help. In general, systematic studies of specific phobias are lacking in the older population.

OBSESSIVE-COMPULSIVE DISORDER

Obsessive-compulsive disorder (OCD) involves persistent patterns of thoughts, obsessions and behaviors, compulsions that are performed in an effort to decrease the anxiety experienced as a result of the thoughts. Obsessions are thoughts or ideas that come to a person's mind, frequently during the process of completing a specific task, or that occur during a particular type of situation. For example, sufferers may find themselves washing their hands repeatedly, for hours at a time, as a result of shaking a stranger's hand. The unwanted thought is that they may have exposed themselves to a serious disease. The act of washing in this example is what is referred to as a compulsion. OCD is a disorder that is chronic and often disabling for the individual¹⁶. Depression and other symptoms of anxiety often accompany the symptoms of OCD.

POST-TRAUMATIC STRESS DISORDER (PTSD)

The distinctive feature of post-traumatic stress disorder (PTSD) is that the individual has experienced, either witnessed or was a victim of, a traumatic event to which they reacted with feelings of fear and helplessness. Examples of such events include those that involve actual or threatened death or serious injury, or other threat to one's integrity, or witnessing an event that involves death or serious injury of another, or hearing about death or serious injury to a family member or close associate. In addition, the individual's accompanying response must have involved extreme fear, helplessness or horror. Other essential features include a number of symptoms that cluster into three categories: (a) persistent *re-experiencing* of the traumatic event; (b) persistent *avoidance of stimuli* associated with the traumatic event and a numbing of general responsiveness; and (c) persistent symptoms of *increased arousal*. Symptoms of re-experiencing include distressing dreams of the event, and intense physical and/or psychological distress at exposure to internal or external cues that

symbolize or resemble an aspect of the event. Symptoms of avoidance and numbing include efforts to avoid activities, people, places, and conversations associated with, or that would arouse recollections of, the trauma. Symptoms of hyperarousal include difficulty in falling or staying asleep, hypervigilance and exaggerated startle response. PTSD usually presents with co-morbid conditions such as depression, panic disorder and substance use disorders. Symptoms must be present for at least 1 month and cause clinically significant distress or impairment in social, occupational or other important area of functioning. Reports of PTSD in elderly Holocaust survivors¹⁷ and among elders who were prisoners of war during World War II¹⁸ indicate that PTSD can be a chronic disorder, continuing into old age. There is some evidence that the intensity of the physiological response to the original trauma may be the most significant predictor of a relatively poor outcome and a chronic course¹⁹. It also appears that ongoing life stressors may slow the recovery process.

ACUTE STRESS DISORDER (ASD)

Characteristic features of this disorder include the development of anxiety, dissociative and other symptoms that occur between 2 days and 1 month after exposure to an extreme traumatic stressor (such as natural or man-made disasters, rape, combat, assault). The symptoms are identical to those described in PTSD, therefore one should consider PTSD as the diagnostic descriptor after a month has passed.

GENERALIZED ANXIETY DISORDER

The distinctive symptoms of this disorder include intense worry about more than one area of one's life. This concern is accompanied by symptoms including: feeling easily tired, experiencing other physical symptoms, such as muscle tension, having trouble sleeping through the night, difficulty concentrating on a task, and feeling irritable or on edge. These symptoms need to be described as having occurred for at least 6 months and must be accompanied by the sense that one cannot control the feelings of anxiety. Many elderly patients with this syndrome may also present with features of depression, thus making it difficult to distinguish between the two diagnoses.

ANXIETY DISORDER DUE TO A GENERAL MEDICAL CONDITION/SUBSTANCE-INDUCED ANXIETY

The elderly as a group are probably most prone to developing this syndrome due to their high prevalence of medical illness and the relatively common occurrence of polypharmacy. Generalized anxiety and/or panic symptoms are the usual presentations among these patients. Among the more common medical disorders producing symptoms of anxiety are endocrine conditions (e.g. hyper- and hypothyroidism, hypoglycemia), cardiovascular conditions (e.g. congestive heart failure, pulmonary embolism, angina, arrhythmias), pulmonary conditions (e.g. chronic obstructive pulmonary disease, pneumonia) and neurological conditions (e.g. neoplasms, Parkinson's disease). Among the more common substances/medications producing symptoms of anxiety in the elderly are alcohol (intoxication or withdrawal), stimulants (caffeine, sympathomimetics in over-the-counter medications), steroids, thyroid preparations, anticholinergic medications and antidepressants. A thorough history, with an attempt to clarify temporal relationship of symptomatology with

the onset of medical illness or the beginning of medication, goes a long way toward resolving the issue.

ANXIETY DISORDER NOT OTHERWISE SPECIFIED

This category includes disorders with prominent anxiety symptoms or phobic avoidance behaviors that do not meet criteria for any specific Anxiety or Adjustment Disorder with anxiety features.

MIXED ANXIETY DEPRESSIVE DISORDER (CATEGORY TARGETED FOR FURTHER STUDY IN DSM-IV)

This category of symptoms is included in the DSM-IV Criteria for Further Study. The essential feature of this proposed disorder is dysphoric mood that has been present for at least 1 month. This mood state must be associated with a minimum of four additional symptoms, such as irritability, worry, sleep disturbance, anticipating the worst, concentration or memory difficulties, and hopelessness. Clinicians working with the elderly have long observed the significant overlap in symptoms of anxiety and depression. In fact, it is quite common to see individuals with a combination of anxiety and depression, although one or both disorders might only be present at subsyndromal levels. Since the distinction between symptoms of anxiety and depression may be particularly difficult to make in older populations²⁰, this category has the potential for significant clinical utility. Further, making a distinction between a primary anxiety disorder and depression is not only of theoretical interest but also of considerable pragmatic value, since medications used for these disorders may have very different side-effect profiles.

ANXIETY/AGITATION IN DEMENTIA

Dementia patients, whether living at home or in a long-term care institution, commonly display behaviors described as agitation. Agitation is operationalized as verbal or motor activity that is either appropriate behavior but repeated frequently, or inappropriate behavior that suggests lack of judgment. As many as 85% of dementia patients go on to develop disruptive, agitated behavior. Early identification of triggers, including environmental stimuli, medication side effects and the inability to communicate internal needs, can lead to effective treatment and relief for already overburdened caregivers. One of the unique aspects in treating such patients is the need to also assess the health and function of the caregiver, which is frequently compromised due to the immense stress involved in performing the tasks to keep such patients safe and their needs attended to.

CONCLUSION

In summary, it appears that anxiety disorders are characterized by a chronic course, usually lasting into old age. Most of the research in the area of the phenomenology of anxiety disorders has been carried out in younger populations, and generalization to an older population is only extended by implication and a limited number of clinical studies and not on a broad-base of empirical data. Clinicians should keep in mind certain factors that can make assessment of anxiety in the elderly problematic. These include a higher rate of medical co-morbidity, which can confound the clinical picture in this population. In addition, a mixed symptom picture of anxiety and depression can make accurate assessment

and specific treatment difficult at times. Finally, a tendency to deny psychopathology and a preference for somatic expression of distress may make it difficult at times to accurately assess the extent of anxiety. Studies designed to investigate any differential manifestations of anxiety disorders in the elderly are clearly needed.

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Prognosis of Anxiety Disorders

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Anxiety disorders most commonly begin in early adulthood, seem to have a relatively protracted course and usually continue in later age¹. As expected, relatively scant information exists at present about the prognosis of anxiety in the elderly. Therefore, this chapter will discuss research findings regarding the prognosis of anxiety disorders in the general population and, of necessity, make inferences about outcome in the elderly.

PANIC DISORDER (WITH AND WITHOUT AGORAPHOBIA)

Panic disorder is a common, usually chronic, illness with fluctuating symptomatology, which may be punctuated by periods of partial remission. Age of onset is typically in the mid-20s, but it may also develop in late life^{2,3}. Panic disorder is commonly associated with considerable psychiatric co-morbidity including depression, obsessive-compulsive disorder, post-traumatic stress disorder and social anxiety disorder^{4,5}. Nearly 50% of those with untreated panic disorder develop co-morbid depression and 43% of these have attempted suicide^{6,7}. It is now quite well established that, if untreated, panic disorder may also lead to alcohol abuse⁸, increased risk for suicide⁷ and, in males, higher than average cardiovascular mortality⁹. Many untreated patients also develop multiple avoidance behaviors (agoraphobia), which are likely to produce serious impairments in social and occupational functioning¹⁰.

Panic disorder rarely resolves without medical intervention¹¹. Although both pharmacological and cognitive-behavioral interventions seem to be effective in the short term¹¹, the long-term effect of these treatments on the natural history of panic disorder is less established. Mounting clinical data favor the SSRIs as first-line treatment for patients with panic disorder. Patients should, in general, be treated for a minimum of 1 year. Those who have experienced previous relapses or who have co-morbid conditions should be considered for long-term therapy¹². Patients who have more than two episodes should be maintained indefinitely.

OBSESSIVE-COMPULSIVE DISORDER

Obsessive-compulsive disorder (OCD) is a chronic and often disabling anxiety disorder that is characterized by recurring obsessions and uncontrolled compulsions. It often occurs comorbidly with a number of depressive and anxiety disorders. Persons with obsessive-compulsive disorder often experience significant personal and social morbidity. Additionally, they may have difficulty finishing school, finding and maintaining a job and developing relationships¹³.

Obsessive-compulsive disorder has a chronic course, and although symptoms may fluctuate over time, the disorder rarely resolves spontaneously without treatment^{14,15}. Until relatively recently, many patients were refractory to conventional pharmacotherapy and obsessive-compulsive disorder was traditionally thought to have poor prognosis. However, of late, advances in both pharmacological and behavioral approaches and their combined use have become effective and important approaches in the management of this disorder. Specific psychopharmacological agents, clomipramine (a tricyclic antidepressant) and the selective serotonin reuptake inhibitors (SSRIs), have proved effective in controlled studies^{16,17,18}. Thus, with appropriate diagnosis and treatments, most patients, including the elderly with OCD, will experience benefits and an improved quality of life.

SOCIAL PHOBIA (SOCIAL ANXIETY DISORDER)

Social anxiety disorder, an often-overlooked diagnosis, is characterized by a marked fear of social performance, excessive fear of scrutiny, and fear of acting in a way that will be embarrassing to oneself. Thus, exposure to social or public situations may provoke an anxiety response and be endured under extreme distress. The mean age of onset is 15.5 years and onset after age 25 years is uncommon¹⁹. Epidemiological data suggest that this disorder is chronic and persisting into old age, with most cases remaining untreated²⁰.

Social anxiety disorder is responsive to both pharmacological and psychological interventions and the two modalities appear to have complementary strengths. For example, cognitive-behavioral therapy²¹ and social skills training have proven value. Pharmacological treatment includes the SSRIs, the monoamine oxidase inhibitors and the benzodiazepines, with the SSRIs as first-line treatment^{22,23,24}. Information about long-term prognosis with these treatment strategies is lacking. A limitation of all medications is the substantial rate of relapse observed even after prolonged treatment. However, there appears to be a lower incidence of relapse following discontinuation of CBT²⁵. Systematic studies of this disorder in the elderly are non-existent, although our clinical experience suggests that public speaking may seem less frightening, and a phobia such as fear of eating in public may be more bothersome, to the elderly than to younger people.

SIMPLE PHOBIA (SPECIFIC PHOBIA)

Generally, specific fears and phobias may be classified into the following groups: (a) situational phobias (lightning, enclosed

spaces, darkness, flying, heights); (b) animal phobias (spiders, snakes); (c) blood-injury (injections, dentists, blood, injuries). An analysis of Epidemiological Catchment Area data suggests that the onset of social phobia is associated with female gender, low education and never having been married^{26,27}. It is difficult to characterize the longitudinal course of simple phobias, principally due to the multiplicity of specific causal stimuli. In addition, individuals with one specific phobia may develop additional phobias or other psychopathology at some point in the course of their affliction.

Exposure and related desensitization techniques are the psychosocial treatments of choice for all variants of specific phobias and promise significant improvement²⁸. These strategies also promise effective biobehavioral interventions for older individuals.

POST-TRAUMATIC STRESS DISORDER (PTSD)

This condition is commonly characterized by an acute-on-chronic course of multiple symptoms (e.g. emotional numbing, hyperarousal, hypervigilance, nightmares, avoidance behaviors) after a traumatic event. Most studies suggest that women are not at greater risk for traumatic exposure, but are more likely to develop PTSD when exposed to trauma, especially if experienced prior to age 15 years^{29,30}. Intensity of the physiological response to the original trauma seems to be the most significant predictor of a relatively poor long-term outcome³¹ (see also comments regarding acute stress disorder). Dissociative phenomena, sensation-seeking/high-risk behavior, emotional constriction, and drug and alcohol abuse also seem to indicate poor prognosis; in addition, on-going life stressors may slow the recovery process³¹. PTSD is frequently accompanied by obsessive-compulsive disorder, phobias, dissociative disorder, generalized anxiety disorder, panic disorder, depression and substance use disorders¹³. In addition, a number of somatic symptoms such as headaches, chronic pain, irritable bowel syndrome and fatigue, are commonly co-morbid.

There is accumulating evidence that pharmacotherapy is effective for the treatment of PTSD. For example, the selective serotonin reuptake inhibitors have demonstrated significant broad-spectrum effects in all the PTSD symptom clusters. They may be considered as first-line (preferred) pharmacological agents³². Other medications that may also be considered, are nefazodone, the tricyclic antidepressants and the monoamine oxidase inhibitors. Psychotherapy can be considered as either an alternative or an additive treatment to medications. Numerous psychotherapeutic techniques can help alleviate symptoms. These include cognitive-behavioral therapy, prolonged exposure, supportive-psychoanalytic therapy and stress inoculation training³³.

Symptoms of the disorder are similar across age groups—re-experiencing the trauma, avoidance and hyperarousal—and there is no current evidence that aging affects the development or presentation of PTSD in older individuals. Elderly individuals do not appear any more predisposed to develop PTSD than do younger persons³⁴. It is not uncommon for individuals who have experienced trauma (e.g. combat) to experience an exacerbation of PTSD, or for post-traumatic disorder to be reactivated, during later life^{35,36}. As with other anxiety disorders, pharmacological and biobehavioral interventions found effective with younger populations can be incorporated into treatment for older adults.

GENERALIZED ANXIETY DISORDER (GAD)

GAD typically has an early onset with an acute-on-chronic course and is associated with increased utilization of medical and mental health services and increased consumption of psychotropic

medications³⁷. The presence of a co-morbid diagnosis is associated with a worsened prognosis and reduced remission rates compared with those patients with GAD alone^{38,39}. Women with GAD are more likely to develop co-morbid conditions (e.g. depression) and the presence of such co-morbidity may reduce the likelihood of remission⁴⁰.

Treatments for GAD include both pharmacological and psychological interventions. Efficacy has been reported with buspirone, the benzodiazepines, the SSRIs and venlafaxine^{41,42,43}. Cognitive-behavioral therapy (CBT) can be quite effective for this disorder in the short term⁴⁴. Additionally, the benefits of CBT appear to be maintained at long-term follow-up and thus may provide a long-term and cost-effective intervention for GAD⁴⁵.

ANXIETY DISORDER DUE TO A MEDICAL CONDITION

This syndrome may be more common in the elderly due to more frequent medical illness. Prognosis depends on the nature and course of the underlying medical condition and its management.

ACUTE STRESS DISORDER

Acute stress disorder describes post-traumatic stress reactions that develop in the first month following a traumatic event. A review of the empirical literature on psychological reactions to trauma suggests that dissociative, intrusive, avoidance and arousal symptoms have often been identified across different kinds of traumatic events⁴⁶. Of those individuals who experience trauma, a minority develops acute stress disorder. However, the literature suggests that a substantial majority of those who meet criteria for this disorder subsequently meet the criteria for ASD⁴⁷. Symptoms with strong predictive power for the later development of PTSD include dissociation, re-experiencing, avoidance, acute numbing, and motor restlessness^{48,49}. Therefore, in terms of prognosis, it is important to identify this pattern of reactions and to provide appropriate interventions to minimize their degree and duration.

SUBSTANCE-INDUCED ANXIETY DISORDER

A clinical picture of prominent anxiety, panic attacks, obsessions or compulsions characterizes this disorder. There must be evidence that medication use or substance intoxication or withdrawal are etiologically related to the symptoms. Symptoms must be clearly in excess of those customarily associated with the substance and these must cause clinically significant distress or impairment that warrants independent clinical attention. Once the substance is discontinued, the anxiety symptoms will usually remit within days to several weeks¹³. Symptom resolution is dependent upon the half-life of the substance, the presence of a withdrawal syndrome and other factors such as general health, medical co-morbidities and any psychiatric co-morbidities. For these reasons and the factor of aging, prognosis in the elderly may be more protracted.

CONCLUSION

In summary, it appears that anxiety disorders are characterized by a chronic course, with symptomatology becoming worse during periods of physical and emotional stress. Both pharmacological and psychotherapeutic approaches seem to be effective in the acute or short term. Definitive literature about the longer-term effects of treatments on the natural course of these disorders is still

in progress. Not surprisingly, we know little about the natural course and prognosis of anxiety disorders in old age. Future studies designed to answer these questions are sorely needed.

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Acute Management of Anxiety and Phobias

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Anxiety disorders are among the most common psychiatric conditions occurring in the elderly¹. Although most of these disorders are of rather chronic nature, acute exacerbations, under a variety of conditions, can produce states of extreme anxiety and agitation that can require immediate attention. For example, patients in treatment for an anxiety disorder may seek an emergency appointment with their doctor/therapist due to a magnification of symptoms. Similarly, in a general hospital a psychiatrist may be called to consult on the management of acute anxiety of an elderly patient on a medical or surgical floor; or a resident may be called upon to evaluate and manage an acutely anxious elderly patient in the emergency room. It is also not unusual for patients with a primary anxiety disorder to present for emergency room services, thinking that they have acute medical problems², e.g. patients having a panic attack might fear an impending heart attack and seek medical treatment. Since older anxious patients are more likely to have concomitant medical problems than their younger counterparts, they may require a careful medical evaluation to rule out any organic causes of anxiety³.

The goal of acute management is relief of marked distress; thus, the treatment approach described here will focus on decreasing patients' symptomatology to manageable proportions as expeditiously as possible. Therefore, treatment approaches for long-term management of anxiety and phobias will be omitted. Further, most situations requiring acute management will necessitate combined pharmacological and psychological interventions. Due to a lack of systematic studies of anxiety management in the elderly, much of our discussion will be based on evidence from studies in younger populations and our own clinical experience.

GENERAL PRINCIPLES OF ACUTE MANAGEMENT

Although acute management of various anxiety disorders may vary somewhat according to the diagnosis, certain guidelines can be useful in a majority of situations. To begin, it is important to remember that during states of extreme anxiety, patients can manifest grossly impaired judgment and might appear to be suffering from a psychotic condition, but a few minutes of questioning will usually clarify the issue. A supportive interaction with the patient is essential to successful treatment. A calm, reassuring manner can be very comforting in itself to alleviate the terror of extreme anxiety. Having a keen awareness of the unique psychosocial issues of the elderly, including retirement, possible deaths of close friends and loved ones and a gradual deterioration of physical functioning, is usually very helpful in developing the

Table 102.1 For patients: understanding your anxiety

- The intense physical symptoms you experience when you are highly anxious are those that are natural to the human body; they are not harmful to you as such. All people have an instinctive "fight-or-flight" response to danger. It is the apparent lack of real danger to you that makes your feelings of fear or anxiety seem so uncomfortable and overwhelming.
- A number of factors may have led to your anxiety experience(s). Your doctor or another clinician may have discussed some of these with you. You may have been given medication to help control your anxiety. It is important for you to take your medication exactly as directed. Anxiety may also be controlled by other methods. Some of these are breathing and muscle relaxation skills, visualization (imagination) techniques, and exposure to anxiety-producing situations with the aid of a therapist. No matter what type of treatment you receive, keep your therapist or doctor aware of any problems, questions or concerns you may notice.
- You and your doctor/therapist will be working together to help you in understanding and controlling your anxiety. In your efforts to cope (deal) with anxiety reactions, it is important to keep in mind that you are not alone. Many people suffer from intense and seemingly overwhelming periods of anxiety. Remember also, "there is light at the end of the tunnel". Anxiety symptoms can be controlled. This may take time, practice, courage and a "stick-to-it attitude", but it is definitely do-able.

initial rapport that will allow the patient to comply with subsequent treatment.

Patient education about their condition and various forms of treatment can be especially beneficial to geriatric patients. Patients with an anxiety disorder benefit from a discussion of thoughts, feelings and behaviors with their therapist or doctor, which can enhance rapport and facilitate patient understanding, decrease global anxiety and foster patient compliance. Information that may be provided to patients during acute management of anxiety reactions is included in Table 102.1.

The specific interventions may be pharmacological and/or psychological. Before describing specific treatments for various anxiety syndromes, it will be helpful to review general principles of pharmacological and psychological therapies with older adults.

PHARMACOLOGICAL MANAGEMENT

Common age-related changes in absorption, distribution, protein binding, metabolism and excretion of drugs and their implications have been covered in detail elsewhere in this book, and thus we will only address the relevance of these changes to anxiolytics,

including benzodiazepines, β -blockers and buspirone. For benzodiazepines, the net effect of these changes is usually a relatively higher level of active medication or its metabolites compared to younger people. In addition, an increase in the proportion of body fat with aging may mean that a strongly lipophilic benzodiazepine, such as diazepam, will lead to a much higher accumulation in tissues, compared to less lipophilic drugs such as lorazepam and oxazepam, which are preferable in the elderly⁴. Of the β -blockers, propranolol is the most frequently used. One should note, however, that its adverse side effects are most common in patients over the age of 60, including its potential to cause depression and worsen cardiac failure and bronchial asthma, and its potentially troublesome interactions with various other drugs, such as calcium channel blockers, cimetidine and chlorpromazine⁵. Aging does not seem to significantly alter the pharmacokinetics of buspirone⁶. Appropriate usage of these agents in specific situations will be discussed in later sections.

PSYCHOLOGICAL MANAGEMENT

From a cognitive-behavioral perspective, anxiety can be understood to have three core components: psychological (e.g. cognitions and affects), physiological (e.g. increased heart rate, dizziness) and behavioral (e.g. ruminations, compulsions and avoidance behaviors). When unfounded, severe anxiety initiates and maintains maladaptive functioning and psychological disturbance. How an individual perceives, understands and functions with anxiety can be shaped by such factors as coping mechanisms, personality, social and environmental influences and past trauma. Cognitive-behavioral principles are very effective with a variety of psychiatric symptoms⁷⁻⁸. For example, breathing and muscle relaxation training, guided imagery, systematic desensitization, relabeling of anxiety reactions, insight into irrational beliefs and systematic homework assignments are effective interventions that may be utilized during acute presentations of anxiety. During acute management, elders may need a greater amount of reassurance and doctor/therapist contact time than their younger counterparts. Maintenance of such techniques through follow-up sessions will increase the internal support strategies of the patient and decrease the risk for future crises.

Diagnostic Categories Most Commonly Requiring Management

Panic Disorder

There are several management strategies that have shown some degree of success in panic disorder⁹. The therapeutic efficacy of antidepressants in panic disorder and agoraphobia is quite well established. Particularly effective are the tricyclic antidepressant imipramine, the monoamine oxidase inhibitor (MAOI) phenelzine, and the selective serotonin reuptake inhibitor (SSRI) sertraline¹⁰⁻¹². Due to their rapid onset of therapeutic action, however, benzodiazepines should be considered the mainstay of acute management, as antidepressants usually take approximately 2-3 weeks for their therapeutic effects to take place. Although alprazolam is the most commonly used benzodiazepine in panic disorder¹³, clonazepam¹⁴ and lorazepam¹³ have reportedly been effective.

Cognitive and behavioral therapies are inextricably intertwined in the acute treatment of panic disorder, with or without agoraphobia. Panic disorder may be managed acutely with breathing and muscle relaxation techniques, examination of cognitive beliefs and a series of progressive behavioral exercises. With therapist-assisted graded exposure beginning even during

the acute management phase of treatment, frequent exposure sessions may facilitate the lessening of the anxiety symptoms.

Social Phobia (Fear of Public Speaking, Eating in Public, etc.)

β -Blockers have been shown to be superior to a placebo for treatment of a fear of public speaking or performance anxiety in the general population^{15,16}. The espoused mechanism of such therapeutic response is the suppression of peripheral responses of anxiety (e.g. palpitations). We know of no studies or clinical reports that address the effectiveness of β -blockers in the socially phobic elderly; thus, it is hard to say whether this treatment will be equally effective in the elderly. It is also not clear whether a benzodiazepine in low dose (e.g. 0.5 mg lorazepam) will be helpful in encountering the phobic situations.

Office-based social skills training as well as exposure *in vivo* (individual or group treatment) can be very effective¹⁷. For acute management purposes, teaching a single, generally acceptable "coping strategy" is most useful to patients and can be implemented quite easily in a variety of situations. A skilled clinician may also consider the use of paradoxical intention, visualization and systematic desensitization in acute management interventions.

Specific Phobia (Crime, Medical and Dental Procedures, etc.)

A minority of individuals seek psychiatric treatment for simple phobias, and clinically significant improvement is usually obtained in 75-85% of specific phobias treated¹⁸. Common fears of older adults include being a crime victim and fears about medical and dental procedures. Although crime rates decrease with age, medical and dental procedures increase, therefore successful management strategies are warranted. We find that low-dose benzodiazepines before the medical or dental procedure in very fearful patients may be helpful in alleviating anxiety and producing better compliance with treatment.

Acute management of most simple phobias can be treated effectively, and with therapy gains maintained, with one (2-3 h) office-based, therapist-assisted exposure session¹⁸. Effective treatment requires focusing on one phobia-related avoidance behavior per session. Breathing and muscle relaxation techniques can also be quite effective in suppressing anxiety responses in older adults.

Generalized Anxiety

In certain instances, the symptomatology of patients with a generalized anxiety disorder can become extremely severe and may require immediate intervention with benzodiazepines. We recommend replacing benzodiazepines with buspirone and/or cognitive-behavioral interventions, including the range of relaxation exercises, until the acute symptomatology is under control.

CONCLUSION

Principles for the acute management of anxiety in the elderly remain more or less consistent over the range of anxiety disorders, although the contexts in which one is asked to evaluate and manage such cases may vary greatly. The importance of a good history, empathy to the patient's psychosocial situation, and awareness of a possibility of an underlying medical condition cannot be overemphasized. Finally, one needs to be cognizant of

the great individual variation in this group and should be ready and willing to tailor the usage of medications and/or cognitive-behavioral techniques to each patient's special needs.

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Psychopharmacological Treatment of Anxiety

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Even though the prevalence of anxiety disorders declines as people age, anxiety disorders still remain the most common psychiatric illness in the elderly¹. This statistic represents not only the continuation of chronic anxiety disorders into later life for many people, but also the development of new anxiety disorders for others. Loneliness and fear of isolation, diminished sensory and general functional capacities, increased incidence of illness, financial limitations and the prospect of dying often generate considerable anxiety². Elderly patients who recover from a mood disorder often develop persistent anxiety, especially in the morning. An estimated 10–20% of older patients experience clinically significant symptoms of anxiety³. Unfortunately, many individuals may never seek treatment, or their anxiety may not be recognized. The result has been a significant undertreatment of a very treatable illness.

This chapter reviews the pharmacologic treatment of anxiety disorders in the elderly. At this time, there is no “perfect” anxiolytic drug for treating the elderly^{4,5}. Further, there are significant gaps in the research of anxiety disorders in the elderly, which makes the choice of treatment difficult⁶. This chapter will therefore begin with a review of general considerations a physician must make prior to selecting and starting pharmacologic treatment. A discussion of current pharmacologic options will follow. Classes of medication, rather than specific recommendations for each anxiety disorder, will be reviewed, since research has not clarified primary treatments for most anxiety disorders. Finally, guidelines for the evaluation and selection of pharmacologic treatments are suggested.

GENERAL CONSIDERATIONS

When to Treat

The decision to treat the anxious older patient with medication depends on the severity of the anxiety and the degree to which it interferes with the patient’s functioning^{7,8}. Anxiety may interfere with social and interpersonal activity in the older patient, resulting in a breakdown of support systems or coping skills. It may worsen cognitive function by decreasing a patient’s concentration. Anxiety may also exacerbate physical illnesses or may be an unrecognized consequence of a medical disorder. Anxiety has been related to blood pressure variability and, by extension, to increased cardiovascular risk⁹. The DSM-IV has identified several subtypes of anxiety disorders (Table 103.1) for the general adult population. They are based on the presence of a cluster of symptoms with a characteristic course and treatment. However, anxiety may also present as a symptom of another disorder.

Therefore, the first task is to assess the impact of the anxiety symptoms on social and emotional functioning or the severity of a coexisting physical illness.

Differential Diagnosis

The diagnosis of anxiety, as either a disorder or a symptom, is not always apparent. This is especially true in the elderly patient. Elderly patients are often less willing to discuss “anxiety”, but may report “anxiety-equivalent” complaints and physical illnesses. Thus a patient may deny being “anxious”, but admit to being “jittery”, “sick”, “uneasy”, “flustered”, “hot”, “restless”, “ill”, “achy”, “agitated” or “bad”. Alternatively, the patient may verbalize physical symptoms, such as being “sick to my stomach”, or having “heart pain” or “insomnia”. These complaints may obscure the true diagnosis or complicate another. The physician must therefore be attuned to what the patient is actually saying.

Further complicating the differential diagnosis is the fact that anxiety may present as a primary disorder (panic disorder, generalized anxiety disorder, obsessive-compulsive disorder, etc.) or a symptom of another primary diagnosis (such as depression, thyroid disease, cardiovascular disease or dementia). There are many conditions that may cause anxiety as a symptom (see Table 103.2); thus, differentiating the source or sources may be difficult. Common medical disorders associated with anxiety as a symptom in the elderly include chronic obstructive pulmonary disease, coronary artery disease, early dementia, major depressive episodes, and medication interactions or withdrawal. Environmental stressors, bereavement or anniversary reactions and experience of medical illness or hospitalizations are also commonly associated with anxiety in late life¹⁰. Non-prescription medications (especially caffeine-containing products, certain cold

Table 103.1 DSM-IV anxiety disorders

Panic disorder without agoraphobia
Panic disorder with agoraphobia
Agoraphobia without a history of panic disorder
Specific phobia
Social phobia
Obsessive-compulsive disorder (OCD)
Post-traumatic stress disorder (PTSD)
Acute stress disorder
Generalized anxiety disorder (GAD)
Anxiety disorder due to . . . [a general medical condition]
Substance-induced anxiety disorder
Anxiety disorder, NOS

Table 103.2 Medical disorders associated with anxiety as a symptom

Cardiopulmonary
Asthma
Chronic obstructive pulmonary disease
Hypoxic states
Angina pectoris
Mitral valve prolapse
Cardiac arrhythmias
Congestive heart failure
Cerebral arteriosclerosis
Hypertension
Pulmonary embolism
Neurologic
Partial complex seizures
Early dementia
Delirium
Post-concussion syndrome
Cerebral neoplasm
Huntington's disease
Multiple sclerosis
Vestibular dysfunction
Endocrine
Carcinoid syndrome
Cushing's syndrome
Hypoglycemia; hyperinsulinism
Hypo- or hyperthyroidism
Hypo- or hyperparathyroidism
Menopause
Pheochromocytoma
Premenstrual syndrome
Medications
Anticholinergic medications
Caffeine
Cocaine
Steroids
Sympathomimetics
Alcohol
Narcotics
Sedative-hypnotics

and flu medications, alcohol or nicotine withdrawal, and certain herbal remedies) may contribute to anxiety symptoms.

Adult Studies

Research in the treatment of anxiety disorders for elderly patients is limited. A recent summary of the National Institute of Mental Health Workshop on Late-life Anxiety⁶ has highlighted this problem, noting three significant research gaps: (a) little consensus on the "best" approach to measure and count anxiety symptoms, syndromes or disorders in late life; (b) insufficient numbers of studies that examine anxiety among older adults; and (c) limited knowledge of the differences in "early" and "later" onset of various anxiety disorders. These limitations become especially significant in treatment recommendations for elderly patients with anxiety. A recent review of the literature⁵ indicated that there are very few controlled clinical trials of medication or psychosocial interventions for anxiety disorders in the elderly. Many of the findings take the form of case reports, case series or open studies. Therefore, treatment decisions for elderly patients are usually extrapolated from the clinical studies of younger mixed-age adult populations and personal clinical experience. For the most part, there is little reason to doubt the applicability of the studies to the

elderly patient, yet the clinician should be aware of the limitations of the research, and sensitive to the developing research in this area.

Special Adaptations for the Geriatric Patient

Before prescribing anti-anxiety agents for the elderly, the physician should be aware of the several age-related physiologic changes that may alter drug pharmacokinetics and contribute to increased risk of adverse reactions. These include changes in drug absorption, drug distribution, protein binding, cardiac output, hepatic metabolism and renal clearance¹¹⁻¹³. In addition, changes in neurotransmitter and receptor function in the central nervous system (CNS) may make a patient more sensitive to psychotropic drugs¹⁴. In general, the usual starting dose of psychotropic drugs for geriatric patients is roughly one-half of the starting dose for younger adult patients.

PSYCHOPHARMACOLOGIC DRUGS

During the past three decades, a variety of agents have been used for the treatment of anxiety and anxiety disorders with varying degrees of success. These include benzodiazepines, buspirone, tricyclic antidepressants (TCAs), monoamine oxidase inhibitors (MAOIs), serotonin selective reuptake inhibitors (SSRIs), newer mixed-action antidepressants, antipsychotic neuroleptics, β -blockers and antihistamines. Despite the multiple medications available, none are completely safe or completely satisfactory in the treatment of anxiety. Zimmer and Gershon's¹⁵ conclusion that the "ideal geriatric anxiolytic" has yet to be developed still holds true today. Therefore, effective methods of treating anxiety disorders are especially dependent upon thoughtful, comprehensive and accurate assessment of psychiatric, social and medical status, as well as a thorough knowledge of the patient's drug history and medication options.

BENZODIAZEPINES

Since the 1960s, the benzodiazepine class of compounds has been the mainstay of drug treatment for patients with situational anxiety, GAD and panic disorder¹⁶. They are also frequently prescribed for other indications, such as insomnia, relaxation prior to certain medical procedures, seizures, or agitation in demented patients. In the last two decades, increased attention has been given to the prescription pattern of benzodiazepines in the elderly. Benzodiazepines were prescribed at a much higher rate among elderly patients than in the general population¹⁷. Epidemiologic data suggest that benzodiazepines may be overused in the general elderly population¹⁸. This is significant because of the potential toxicity and side effects of benzodiazepines, especially common in the elderly.

Despite their multiple uses, benzodiazepines are usually classified in two groups: anxiolytics and sedative-hypnotics. Currently, seven benzodiazepines are available for the treatment of anxiety. Listed in their order of introduction, they are chlordiazepoxide, diazepam, oxazepam, clorazepate, lorazepam, alprazolam and clonazepam¹⁹. Prazepam and halazepam, two benzodiazepines previously used for the treatment of anxiety, are no longer available in the USA. The most common sedative-hypnotics are triazolam, temazepam and flurazepam. Commonly used benzodiazepines in the elderly are listed in Table 103.3.

Table 103.3 Commonly used benzodiazepines in the elderly

Generic name	Trade name	Onset of action	Indication	Half-life (h)	Metabolism	Active metabolites	Geriatric dose (mg/day)	Route of administration
Short to intermediate half-life								
Triazolam	Halcion	Fast	Hypnotic	2–5	Conjugation	None	0.125–0.5	Oral
Oxazepam	Serax	Intermediate to slow	Anxiolytic	5–15	Conjugation	None	5–30	Oral
Alprazolam	Xanax	Intermediate	Anxiolytic	6–15	Oxidation	Yes	0.125–3.0	Oral
Lorazepam	Ativan	Intermediate	Anxiolytic	10–20	Conjugation	None	0.5–3	Oral, IV, IM
Temazepam	Restoril	Intermediate to slow	Hypnotic	12–24	Conjugation	None	15–30	Oral
Long half-life								
Chlordiazepoxide	Librium	Intermediate	Anxiolytic	8–30	Oxidation	Yes	5–30	Oral, IV, IM
Diazepam	Valium	Fastest	Anxiolytic	26–53	Oxidation	Yes	2–10	Oral, IV, IM
Clorazepate	Tranxene	Fast	Anxiolytic	30–200	Oxidation	Yes	7.5–15	Oral
Flurazepam	Dalmane	Fast	Hypnotic	64–150	Oxidation	Yes	15	Oral
Klonopin	Clonazepam	Intermediate	Anxiolytic	30–40	Oxidation	Yes	0.25–3.0	Oral

Pharmacokinetics

Benzodiazepines undergo two kinds of biotransformation: oxidation and glucuronide conjugation. Oxidative transformation occurs slowly, giving the drugs a long half-life and producing many active metabolites. Conjugative transformation differs from oxidative transformation in that it occurs rapidly and the metabolic products are pharmacologically inactive. As a general rule, benzodiazepines that are inactivated by conjugation reactions appear to be less likely to interact with other medications. For example, cimetidine has been found to inhibit the metabolism of benzodiazepines that require oxidation, but not benzodiazepines inactivated by conjugation (such as lorazepam or oxazepam). Table 103.4 lists several important drug interactions with benzodiazepines.

Elimination half-lives of benzodiazepines are variable in the elderly. Usually the effects of ultrashort, short and intermediate half-life benzodiazepines do not carry over to the next day when they are used as a sedative²⁰. Ultrashort benzodiazepines are generally used to treat insomnia rather than daytime anxiety. They can cause rebound insomnia after abrupt discontinuation. Safety concerns about triazolam (i.e. after being noted to cause confusion, agitation and hallucinations) have led to its ban in several European countries¹⁸. Benzodiazepines with long half-lives can significantly contribute to increased risk of falls and hip fracture in elderly patients²¹.

The onset of action after a single dose is primarily dependent upon the drug's absorption rate. Most benzodiazepines are highly

lipophilic. Benzodiazepines that are more lipid-soluble have a faster onset of action because they are absorbed and diffused into central synapses more rapidly^{22,23}. This rapid onset of action can produce euphoria and thereby enhance abuse potential.

Efficacy

Benzodiazepines are effective for generalized anxiety disorder (GAD). Clonazepam has been shown effective for social phobia. Clonazepam and alprazolam are effective in panic disorder. Efficacy and use in post-traumatic stress disorder (PTSD) is limited. A paradoxical reaction has been documented when some patients with PTSD are treated with benzodiazepines. None of the benzodiazepines appear effective for obsessive-compulsive disorder (OCD).

Dependence and Withdrawal

True physiologic dependence, resulting in a withdrawal and abstinence syndrome, develops to benzodiazepines usually after 3–4 months of daily use¹⁴. Withdrawal symptoms are likely to be more severe with abruptly discontinued therapy, and with patients receiving short half-life benzodiazepines, or higher daily doses. The symptoms of withdrawal include tachycardia, orthostasis, intention tremors, diaphoresis, hyper-reflexia, anxiety, insomnia, nightmares, malaise, anorexia, headache, muscle pain and twitching, tinnitus, hyperacusis, photophobia, metallic taste, strange smells and, in more severe cases, hyperthermia, nausea, vomiting, delirium, seizures and psychosis.

Side Effects

Although benzodiazepines have been shown to be effective in younger populations, systematic studies in the elderly are lacking. Judicious use is therefore important, since they may have several significant side effects. Adverse drug reactions to the benzodiazepines are almost twice as common in patients over the age of 70 years compared with those aged 40 years or less²⁴.

Benzodiazepines do tend to produce greater effects on the central nervous system of the elderly than in younger patients^{12,13}. This is due partly to increased target-organ sensitivity and partly to altered pharmacokinetics in the elderly (i.e. duration of half-life and peak blood level)⁷. The most common benzodiazepine side effect is a dose-dependent CNS depression. Symptoms include

Table 103.4 Drug interactions with the benzodiazepines

Drug	Effect
Alcohol	Increased CNS sedation
Neuroleptics	Increased CNS sedation
Narcotics	Increased CNS sedation
Antihistamines	Increased CNS sedation
MAO Inhibitors	Increased CNS sedation
Cimetidine	Increased elimination half-life and decreased clearance of alprazolam, diazepam and chlordiazepoxide
Isoniazid	Decreased metabolism of diazepam
Rifampin	Increased metabolism of diazepam
Antacids	Decreased absorption of clorazepate and chlordiazepoxide
Digoxin	Increased digoxin levels
Levodopa	Decreased control of parkinsonism by levodopa
Fluvoxamine	Increased levels of alprazolam

fatigue, drowsiness, sedation, muscle weakness, blurred vision, nystagmus, dysarthria, ataxia and impaired psychomotor and cognitive performance^{12,25,26}. The impairment of motor coordination causes drivers taking benzodiazepines to be five times more likely of being involved in a serious road accident²⁷. Cognitive impairment can be severe enough to present as a pseudodementia in susceptible elderly patients.

Benzodiazepines may also cause a paradoxical reaction of restlessness, confusion, irritability and even aggression. Outbursts of anger in elderly patients receiving benzodiazepines may indicate the need to consider an alternative medicine. There are also published case reports of benzodiazepines inducing a secondary mania^{28,29}. Benzodiazepines may cause mild respiratory depression in patients with chronic obstructive lung disease. Mixing benzodiazepines with other CNS depressants such as alcohol can lead to severe intoxication or (potentially lethal) respiratory depression.

Selection

In general, benzodiazepines are equivalent in terms of overall efficacy³⁰. The selection of a particular benzodiazepine is primarily based upon the patient's particular problem and the medication properties (route of metabolism, length of half-life, onset of action, and presence of active metabolites)³¹. In general, the following guidelines should be considered when using a benzodiazepine:

1. Benzodiazepines that undergo conjugation to water-soluble glucuronides prior to excretion in the urine (e.g. temazepam, lorazepam, oxazepam) have no active metabolites and their pharmacokinetics are not significantly changed by the aging process³². They are probably the wisest choice for elderly patients with severely impaired hepatic function.
2. Short (but not ultrashort) half-life drugs are preferable to long half-life medications, since they appear less likely to increase the risk for hip fractures²¹.
3. Accumulation of benzodiazepines is directly related to the amount of fat. Therefore, the obese or severely medically frail patients may be at increased risk.
4. Avoid benzodiazepine use in patients dependent on alcohol or other drugs.
5. Begin with lower doses and titrate upwards gradually ("start low and go slow").
6. Try to limit the length of use to 3–4 months. Taper benzodiazepines over a 4–8 week period.

Buspirone

Buspirone (Buspar) is a novel anti-anxiety agent unrelated to the benzodiazepines in chemical structure or pharmacologic characteristics. Its mechanism of action is probably related to its high affinity for the serotonin type 5-HT_{1A} receptor, which causes reduced serotonergic activity. In addition, it enhances brain dopaminergic and noradrenergic activity^{33,34}.

Efficacy

Buspirone is effective in the treatment of generalized anxiety disorder in the elderly. It is well tolerated and as effective as the benzodiazepines^{35–38}. However, buspirone does not appear to be effective in the treatment of panic disorder³⁹. Some researchers suggest that buspirone may be helpful in mixed anxiety/depression

symptoms. It may also be effective as an adjunct treatment for OCD. Its use in PTSD and social phobia appears to be limited.

Administration

Therapeutic doses are in the range 20–60 mg daily; however, buspirone's short half-life (averaging 2–3 h) requires that it be given three times a day (usually with meals). Studies have demonstrated that buspirone may remain effective for at least 6 weeks, although longer efficacy is presumed.

Two major disadvantages of buspirone are the requirement for multiple daily dosing and the lack of immediate effect. Buspirone may take 1–3 weeks at therapeutic dosing before the anxiolytic effect begins. Some researchers have also suggested that the efficacy of buspirone may be reduced in patients who have previously been treated with benzodiazepines⁴⁰. Others suggest using a benzodiazepine for the first 1–2 weeks when initiating treatment with buspirone until it becomes effective.

Side Effects

Buspirone side effects include nausea, headache, nervousness, dizziness, lightheadedness and fatigue. Unlike the benzodiazepines, buspirone does not appear to cause psychomotor impairment, dependence, withdrawal or abuse⁴¹. Further, it does not interact with alcohol and other sedative drugs. Buspirone lacks hypnotic, anticonvulsant and muscle relaxant properties. Therefore, it may be of particular value in the treatment of patients unable to tolerate the sedative effects of benzodiazepines⁴², or patients with a history of substance abuse.

Antidepressants

Tricyclic Antidepressants

Efficacy. Tricyclic antidepressants (TCAs) have been shown to be effective in treating mixed anxiety–depression states, panic disorder and generalized anxiety disorder in the elderly^{43–46}. In the general adult population, TCAs are frequently used in PTSD and clomipramine has been approved by the FDA to treat OCD. However, the overall use of TCAs has decreased as other options (especially the SSRIs) have become available. This is primarily due to the significant side effects TCAs have at therapeutic doses that also increase the risk to physically ill patients and potentiates toxicity in overdose. Further, like all other medications used for the treatment of anxiety (except the benzodiazepines), TCAs usually require several weeks to show maximal benefit. Despite their shortcomings, tricyclic antidepressants remain an alternative treatment for GAD⁴⁷.

Side Effects. Common side effects may be mediated by α -adrenergic blockade, anticholinergic effects and antihistaminergic effects. The α -adrenergic blockade of TCAs may cause significant orthostatic hypotension or cardiac conduction irregularities. The elderly are particularly susceptible to injury from orthostatic falls. Patients with complete heart block should not be given TCAs because these medications can cause a prolonged QRS complex. Trazodone, a heterocyclic antidepressant, is sometimes used as a sedative or in the treatment of agitation for demented patients, but the side effects of postural hypotension may limit its use²⁰.

Anticholinergic side effects of TCAs are dry mouth, blurred vision, constipation, urinary retention and confusion or even psychosis. This may be particularly significant in patients with

Alzheimer's disease or other disorders that impair memory. The major antihistaminergic effect is sedation.

Guidelines for Use. Imipramine and amitriptyline have long been established in the adult population for use in various anxiety disorders. However, their tertiary amine structure tends to cause increased anticholinergic, adrenergic and sedative side effects. The secondary amines, nortriptyline and desipramine, are preferred for use in the elderly due to their less intense side effects. In general, anxiety disorders appear to respond at doses lower than those used in mood disorders. A baseline EKG is highly recommended prior to starting therapy, since TCAs can cause a prolonged QRS complex.

Monoamine Oxidase Inhibitors (MAOIs)

Monoamine oxidase inhibitors have been effective in treating mixed anxiety–depression and panic disorder but not pure generalized anxiety^{44,45}. MAOIs are rarely used now because of their potential side effects (especially the drug–diet interactions) and the wider availability of newer antidepressant choices. Phenzelzine and tranylcypromine are the MAOIs of choice for the elderly in the USA⁸. Interestingly, phenzelzine was found to be more effective in the elderly than in younger patients⁴⁴, and just as well tolerated as nortriptyline⁴⁸. The starting dose is 15 mg daily in the morning, increasing by 15 mg every few days to an average dose of 45–60 mg⁴⁹. Moclobemide, a reversible MAOI available outside the USA, is effective in social phobia and panic disorder^{50,51}.

Orthostatic hypotension is the major side effect of MAOIs; however, the major concern is the possibility of an acute hypertensive crisis due to the drug and dietary interactions.

Selective Serotonin Reuptake Inhibitors (SSRIs)

Efficacy. The introduction of SSRIs has transformed the treatment of both depression and anxiety in the adult population. Five SSRIs are now available for use in the USA: fluoxetine, sertraline, paroxetine, fluvoxamine and citalopram. Since their introduction, certain SSRIs have obtained indications for the treatment of not only depression, but also panic disorder, bulimia, OCD and social phobia. They are also used in the treatment of PTSD and GAD. Still, as with the other medications, the treatment of geriatric patients is primarily extrapolated from studies of younger adults. Case reports, open trials and some controlled studies in elderly patients have been completed in the elderly population. Fluoxetine has been shown to be effective in treatment of geriatric depressed patients with agitation²⁰. Fluvoxamine and fluoxetine have been shown to be effective in clinical trials for OCD that have included some elderly patients^{52,53}. Case studies have shown sertraline, paroxetine, venlafaxine, fluoxetine and fluvoxamine effective in social phobias and anxiety^{54–57}.

Side Effects. All SSRIs can cause gastrointestinal effects (nausea and diarrhea), sexual arousal and performance changes, and a decrease in appetite and weight. The nausea and diarrhea are usually dose-dependent and resolve for most patients within the first week of treatment. An increasing concern has been SSRI-associated weight gain, seen with extended use. Paroxetine has been noted to have some anticholinergic activity that may be more apparent in elderly, sensitive patients. A significant potential side effect of SSRIs includes a tendency to stimulate patients, causing tremor and jitteriness.

Guidelines for Use. Since some SSRIs have been associated with “activation” and worsening of anxiety, some clinicians have suggested avoiding them as an initial choice for treatment²⁰. Smith and colleagues⁵⁸ have recommended that SSRIs that have shorter half-lives and are less activating, such as paroxetine or sertraline, be used preferentially to fluoxetine. However, it should be noted that fluoxetine has been shown to be effective in treating anxiety symptoms associated with depression. Because of the interaction between cytochrome P-450 and drug metabolism, caution should be used when combining SSRIs with tricyclic antidepressants, antiarrhythmics, codeine, carbamazepine, benzodiazepines, and β -blockers or calcium channel blockers.

Newer Antidepressants

Nefazodone offers promise as a useful antidepressant for depressed elderly patients who have concomitant anxiety symptoms²⁰. It has an acceptable level of daytime sedation for the elderly⁵⁹, but minimal anticholinergic and other side effects. Its more moderate serotonin reuptake inhibition may make it less likely to create agitation than the SSRIs⁶⁰. Nefazodone does inhibit P-450 isoenzymes and is contraindicated for use with terfenadine, astemizole and cisapride. It will also increase the plasma levels of alprazolam, midazolam and triazolam.

Venlafaxine, a newer antidepressant that enhances both norepinephrine and serotonin activity, is increasingly being used to treat depression and anxiety in the general adult population⁶¹. It is approved for the treatment of GAD. Its side-effect profile is similar to the SSRIs. Small elevations in blood pressure may be seen at dosages above 200 mg/day.

Mirtazapine is the first of a new class of antidepressants, the noradrenergic and specific serotonergic antidepressants (NaSSA). In trials for antidepressant treatment in the general adult population, mirtazapine has shown beneficial effects on the concomitant symptoms of anxiety and sleep disturbances⁶². It has few anticholinergic, adrenergic and serotonin-related adverse effects, but can be very sedating due to the antihistaminergic effects. There is no current data on its effectiveness in the elderly.

Neuroleptics

Neuroleptics are often useful for treating severe agitation associated with psychosis, delirium, and dementia⁶³. The newer antipsychotics such as olanzapine and risperidone are especially being used in the treatment of refractory anxiety in the context of dementia. However, the use of neuroleptics for the treatment of subjective anxiety states, especially in the elderly, has never been demonstrated⁷. It is important to remember that neuroleptic drug side effects, such as sedation, extrapyramidal reactions, orthostatic hypotension, anticholinergic effects and tardive dyskinesia, can have potentially devastating consequences in older people. Even the newer antipsychotic agents have significant side-effect profiles. Therefore, neuroleptics should not play a significant role in the treatment of anxiety disorders in the elderly.

β -Blockers

The usefulness of β -blockers for treatment of anxiety in the elderly is unclear, since the data for the elderly are restricted to clinical case reports³¹. β -Blocking agents have been shown to be specifically beneficial in younger patients with predominantly somatic symptoms associated with generalized anxiety or anxiety related to stressful situations⁶⁴. They may also have potential use in the treatment of aggression and agitation in patients with

organic brain disease⁶⁵. Although β -blockers decrease autonomic-mediated symptoms such as diaphoresis, palpitation, tremor and gastrointestinal upset, they usually do not reduce the inner subjective effects^{2,4}.

Propranolol in small doses (e.g. 5–10 mg one to four times a day) may be effective in elderly patients²⁰. These drugs should not be used in patients with chronic obstructive pulmonary disease, congestive heart failure, heart block, insulin-dependent diabetes, severe renal disease or peripheral vascular disease.

Antihistamines

Sedating antihistamines such as hydroxyzine and diphenhydramine hydrochloride are sometimes useful for anxiety or insomnia in the elderly. They have been rarely recommended because they are less effective than benzodiazepines, and their anticholinergic side effects are outweighed by their weak anxiolytic effects⁴. They may be used in patients with mild symptoms, in severe chronic obstructive pulmonary disease, addiction-prone personalities, alcoholics, or patients for whom more traditional drugs are not effective⁶⁶. However, physicians must be aware that the elderly patient is much more susceptible to their anticholinergic properties, which may cause blurred vision, tachycardia, dry mouth, urinary urgency, constipation, restlessness, hallucinations and confusion. Antihistamines have no potential for inducing drug dependency or addiction.

GENERAL GUIDELINES

Despite the limited data on treatment of anxiety disorders in the elderly, the clinician can successfully treat patients with a conservative and thoughtful use of medications. The following guidelines have been adapted from Small²⁰.

1. Conduct a complete psychiatric evaluation. Listen specifically for expression of anxiety. Does this patient have anxiety that significantly affects their quality of life or functioning?
2. Consider the full differential diagnosis. Does the pattern of anxiety identify itself as a formal anxiety disorder, or as a symptom of another psychiatric or medical disorder? Geriatric psychiatry has been called “the specialty of co-morbidity”. There may be several potential etiologies for anxiety symptoms that should be considered before initiating treatment.
3. Consider non-pharmacologic treatments first. Education and reassurance are invaluable in the treatment of anxiety, and may themselves be adequate treatments. Specifically address social stressors and evaluate the effectiveness of the support systems. Attention to family caregivers may facilitate the positive response to other treatment. Remember, the ability to benefit from therapy is not based on age.
4. Minimize polypharmacy. In the geriatric population (especially those in the nursing care facilities), the use of multiple medications is the rule rather than the exception. Most clinicians stress the importance of reviewing the medication list for potential areas of reduction, prior to adding new treatments. Reducing the number of medications may actually treat the anxiety symptoms²⁰.
5. When selecting an anxiolytic, consider the full presentation rather than just the anxiety when selecting an initial medication. For example, use an antidepressant if depressive symptoms are apparent. Avoid anticholinergic medications in patients with dementia. Avoid benzodiazepines when the patient’s ability to ambulate is compromised.

6. As far as possible, make medication changes one at a time in order to clarify whether a complaint results from a medication side effect or an underlying illness.
7. “Start low and go slow”.

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Obsessive–compulsive Disorder

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Although obsessive–compulsive disorder (OCD) is known to occur in old age, studies are few and information is limited. This may be because it is often not perceived as a disorder of late life. The mean age of onset is 20–25 years^{1–3} and it is unusual for OCD to have its first onset after the age of 50 years. However, OCD is a chronic disorder if untreated, and a significant proportion of cases persist into old age, when they may present to services for the first time. It is important, therefore, that old age psychiatrists are aware of this disorder and of its management.

CLINICAL FEATURES

Diagnostic Criteria

OCD is characterized by intrusive, persistent obsessive thoughts, images or impulses and/or compulsive behaviours that are a significant source of distress, or interfere with the patient's personal or social functioning. The current diagnostic criteria, as set out in DSM-IV⁴ and ICD-10⁵, apply to all patients, irrespective of age. The limited evidence available indicates that the clinical features of OCD in elderly patients are very similar to those of younger adults. In their comparative study, Kohn *et al.*⁶ found that concerns about symmetry, need-to-know and counting rituals were less common in elderly patients, and hand-washing and fear of having sinned were more common, but otherwise there were few differences in clinical features compared with younger OCD patients. Extreme ego-syntonic religiosity has been proposed as a variant of OCD that may be more common in older patients⁷.

Differential Diagnosis

Unpleasant, intrusive thoughts and abnormal stereotyped behaviours occur in other mental disorders, and OCD is not diagnosed if their content is *exclusively* related to another disorder, e.g. guilty preoccupations in depression, worries in generalized anxiety, concern with illness in hypochondriasis, weight control in anorexia or avoidance in phobic disorders⁴. It should be borne in mind that conditions such as depression, generalized anxiety and substance abuse may be co-morbid with OCD. In elderly patients, increased anxious orderliness may be a prodrome of dementia; however, this behaviour is not resisted or associated with the tension that occurs in OCD. The compulsive behaviours of OCD resemble the stereotyped behaviours that occur in certain other disorders, such as Tourette's syndrome, Sydenham's chorea, encephalitis and partial complex seizures. Tourette's syndrome

and OCD commonly co-occur⁸, and patients with OCD may have a history of Sydenham's chorea in childhood⁹.

Despite its similar name, obsessional personality disorder is quite distinct from OCD. It is characterized not by obsessions and compulsions, but by a preoccupation with orderliness, perfection and control dating back to early adulthood⁴. In some individuals, there is an inability to discard personal possessions, which may present as the so-called "senile squalor" syndrome after a lifetime of accumulated rubbish.

Not all patients with OCD have insight into the irrationality and inappropriateness of their obsessions and compulsions. If the obsessional thoughts are held with delusional intensity, an additional diagnosis of delusional disorder may be warranted. The ruminative delusions and stereotypies of schizophrenia are usually not ego-dystonic, and therefore would not be regarded as OCD⁴.

Clinical Assessment

An effective treatment plan for OCD requires a detailed clinical assessment. What exactly are the main problems? What, if anything, exacerbates or improves the symptoms? How long has the condition been present, and how has it evolved since its onset? What treatments, if any, have been tried in the past? What other symptoms or disorders are present? Any concomitant depression, mania, psychosis or alcohol dependency will require specific management before behavioural treatments for OCD can be effective. If the patient is cognitively impaired, this will have implications for the choice of treatment; for example, some behavioural strategies will not work if information cannot be retained or recalled. In elderly patients with OCD of recent onset, it is important to investigate carefully for any underlying cerebral disease. Late-onset cases are associated with frontal dysfunction¹⁰, which may be caused by a variety of focal and generalized disorders, including cerebrovascular disease, tumours and primary neurodegenerative dementias. Late-onset OCD may also be the result of external factors, such as adverse life events and exposure to trauma, that weaken an elderly individual's resistance to long-standing subclinical obsessional¹¹.

EPIDEMIOLOGY

Most of our knowledge about the epidemiology of OCD in old age derives from the US National Institute for Mental Health (NIMH) Epidemiologic Catchment Area (ECA) Program. Overall, the 1 year prevalence for those aged 65 years and older was

0.85% (men 0.75%, women 0.93%), as opposed to 1.65% for the sample as a whole¹². A more detailed analysis of the elderly population at the Eastern Baltimore site found prevalence rates of 1.3% for those aged 65–74 years and 0.6% in those aged 75+¹³. Following the second wave of the ECA, annual incidence rates were estimated. In males aged 65+, the incidence rate of OCD was one-third of that for males of all ages, but in females there was a non-significant upturn in the incidence rate after age 65¹⁴. In common with a number of other psychiatric disorders, the lifetime prevalence of OCD decreased with age in this study. The reason for this is unclear, but it may be the result of cohort effects, differential mortality or age-specific differences in symptom ascertainment and recall.

AETIOLOGY

Genetic factors play an important role in the aetiology of OCD; it occurs in 40–50% of parents, 19–39% of siblings and 16% of children of probands with the disorder¹⁵. Just what is inherited is not clear; other anxiety disorders are also more common in the families of OCD probands¹⁶. Most of the evidence from clinical, neuropsychological and neuroimaging studies implicates the basal ganglia and their connections with the thalamus and the cerebral cortex in the aetiology of OCD¹⁷. Specifically, it has been proposed that there is dysfunction in a neuronal circuit involving the orbitofrontal cortex, the basal ganglia, the substantia nigra and the ventrolateral pallidum¹⁸. The specific response of OCD to serotonin (5-HT) reuptake-inhibiting drugs (see below) suggests that serotonergic neuronal systems are involved, directly or indirectly. In the cognitive-behavioural model of OCD, obsessions and compulsions result from pathological, anxiety-provoking over-control of normal intrusive cognitions¹⁹.

TREATMENT

There are no randomized, controlled trials of treatment of OCD in elderly patients. Accordingly, the guidelines that follow are based upon case reports and extrapolations from studies in younger adults²⁰.

Non-pharmacological Treatments

Behavioural therapy, in the form of exposure and response-prevention (ERP), is well described as an effective intervention for OCD in younger adults²¹. This involves exposing the patient to the feared situation, and helping him/her to resist the urge to perform the compulsive behaviours that would normally follow this exposure²². ERP is least effective in those who have obsessional thoughts and covert rituals unaccompanied by compulsive behaviour. In these patients, a cognitive approach directed at modifying the misinterpretation of intrusive thoughts is more appropriate²³. There have been a number of case-reports of effective behavioural interventions with elderly OCD patients^{19,24–27}, but most are difficult to interpret because of the concomitant administration of medication. In the case described by Calamari *et al.*²⁷, significant improvement following ERP was maintained without medication at 8 month follow-up.

Freud proposed that obsessional symptoms were a regression to a pregenital anal-sadistic phase of development. However, despite this and subsequent psychodynamic formulations of the disorder, there is no evidence that psychodynamic psychotherapy is an effective treatment for OCD at any age.

Pharmacological Treatments

Studies in younger adults indicate that 30–60% of patients with OCD show improvement on appropriate medication, and that drug treatment appears to be more effective for obsessional thoughts than for compulsive behaviours. In practice, drug treatment and behavioural therapy are often given in combination.

The theory that OCD is a disorder of serotonergic function is based upon the empirical observation that it can be effectively treated by drugs that inhibit serotonin reuptake. Clomipramine is the most extensively studied drug treatment for OCD, and its effectiveness has been established in a number of double-blind placebo-controlled trials in younger adults²⁸. However, the lack of receptor sensitivity means that it has significant anticholinergic and antihistaminergic side effects that limit its usefulness in elderly patients²⁹. In this age group, the drug of first choice is one of the specific serotonin reuptake inhibitors (SSRIs). Fluoxetine, paroxetine and fluvoxamine are currently licensed in the UK for the treatment of OCD, although none of the trials supporting this indication specifically involved elderly patients. The effective dose for the treatment of OCD with these drugs tends to be higher than that required to treat depression, and the time taken to respond is typically much longer: 10–18 weeks. Studies suggest that long-term therapy is required, as discontinuation of medication leads to relapse of symptoms.

There is little evidence for the effectiveness of other drug treatments in OCD. There are some case reports suggesting that monoamine oxidase inhibitors (MAOIs) may be useful in patients with concomitant panic or severe anxiety. Anxiolytic drugs may also help with the anxiety associated with OCD, but do not appear to have any effect on the core symptoms. A possible exception is buspirone, which may augment the effect of fluoxetine^{30,31}. Lithium augmentation of fluoxetine has also been reported as effective in one elderly case³².

Physical Treatments

There is very little evidence to suggest that ECT is effective in the treatment of OCD in patients who are not also depressed³³. Some good results have been reported for stereotactic neurosurgical procedures in patients with severe and treatment-refractory illness, including elderly subjects²⁰, but since negative outcomes are rarely described, this evidence is difficult to interpret.

CONCLUSIONS

OCD may present for the first time in old age, and many elderly patients with chronic illness will not have been exposed to the full range of pharmacological and cognitive-behavioural treatments that are now available. It is important that old age psychiatry services are aware of these treatments and develop some experience in their delivery. There is some evidence that they are effective in elderly patients, but further research is needed. Patients with a new onset of OCD in late life need careful assessment to exclude underlying organic brain disease.

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Hypochondriacal Disorder

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The term “hypochondriasis” has its origins in the ancient Greek language. Anatomically, the hypochondrium refers to that part of the body between the ribs and the xiphoid cartilage. The ancient Greeks believed that this part of the body, especially the spleen, was the seat of morbid anxiety about one’s health, depression, bad mood and simulated disease^{1,2}. This old theory has not withstood the passage of time, but the term “hypochondriasis” has survived and is part of our modern diagnostic nomenclature³⁻⁵. Hypochondriasis has joined the ranks of syndromes known as somatoform disorder in DSM-IV, with the following diagnostic criteria:

1. The predominant disturbance is preoccupation with the fear of having or the belief that one has a serious disease based on the individual’s interpretation of physical signs or sensations as evidence of physical illness (do not include misinterpretation of physical symptoms of panic attack).
2. Appropriate physical evaluation does not support the diagnosis of any physical disorder that can account for the physical signs or sensations or the individual’s unwarranted interpretation of them, and the symptoms in (1) are not only the symptoms of panic attacks.
3. The fear of having, or the belief that one has, a disease persists despite medical reassurance.
4. The duration of the disturbance is at least 6 months.
5. The belief in (1) above is not a fixed delusion, as in delusional disorder, somatic type.

There are issues regarding hypochondriasis as a diagnostic entity. Some clinicians and the *International Classification of Diseases* regard it as a specific non-psychotic psychiatric disorder, while others hold that it is a syndrome (a collection of similar symptoms that occur together but are of multiple etiology). It is evident that hypochondriacal symptoms may be part of another disorder, such as a mood disorder or a defense mechanism, as well as a character trait. Starcevic⁶ has dissected the hypochondriacal syndrome into potentially useful constructs that represent a stepwise progression. The “hypochondriacal core”, as the preoccupation with bodily sensations and functioning, gives rise to a state of “somatic uncertainty”, an insecurity feeling resulting in intense anxiety which is poorly tolerated by the individual. This leads to the “disease suspicion”, which represents the fear of having a disease. “Hypochondriacal behaviors” ensue as the patient obsessively seeks a medical work-up to uncover the cause of his/her symptoms and is generally dissatisfied, as it is found that there is either an absence of physical disease or that the symptoms

are out of proportion to the pathology. Starcevic points out that the hypochondriacal syndrome is a heterogeneous entity with varying degrees of bodily preoccupation, fear, suspicion and a variety of complaints.

Hypochondriasis, with its state of uncertainty, may be seen as a feature of an anxiety disorder. It may be part of a mood disorder, such as a masked depression, where the patient will deny feeling depressed but respond to antidepressant medication. Starcevic⁶ makes the point that hypochondriasis may be incorporated into a long-standing, maladaptive pattern of functioning and that many personality disorders may have a hypochondriacal manifestation. There is a fine line to be crossed when the disease suspicion turns into “disease conviction” and the syndrome is no longer hypochondriasis, but a psychotic disorder, such as paranoid delusional disorder, paranoid schizophrenia or major depression with psychotic features.

In his 1987 review article on the subject matter, Lipowski⁷ considers that predisposing factors such as genetics, developmental learning, personality and sociocultural environment play a role in hypochondriasis. Swedish investigators have gathered data from adoption studies in regard to familial somatization patterns. The subjects of their studies were drawn from 912 women born out of wedlock in Stockholm, Sweden, during 1930–1949. Between 1965 and 1973, the medical records of 859 subjects were reviewed for duration and number of sick leaves, chief complaint and diagnosis⁸⁻¹⁰; 144 were found to be somatizers. This study suggests that somatization is more common in adopted women than in non-adopted women. This raises the issue of genetic predisposition to somatization, as adoptees are known to have a higher percentage of biological parents with alcoholism and criminality compared to non-adoptees. There may be a complex interaction between the type of somatizers, alcoholism and antisocial behavior and sex differences. The interaction between biological predisposition and environmental influences requires additional attention.

Theories that conceptualize the genesis of hypochondriasis to learned behavior from childhood sound intuitively correct¹¹. Children who grow up in families where a serious or chronic illness is present, or who are exposed to a hypochondriacal relative, may well learn a way to obtain attention, sympathy and support, or get the message that physical complaints are acceptable, while complaints of emotional distress are not. Children suffering from physical disorders may get anxious attention from parents. They may also learn that being sick is to avoid unpleasant duties. Somatization may serve as a way

to deal with adverse social situations and to maintain self-esteem.

Kanner² felt that hypochondriacal attitudes in children often reflected school problems or "unhappiness at home". He observed that some mothers focused on their child's somatic functioning rather than their own, thus teaching the child to somatize. This improper maladaptive coping pattern may well continue in adulthood. While it is possible that these early childhood influences shape the child into a hypochondriacal character, it may not necessarily follow. There may be innate personality characteristics that predispose to somatization. Barsky and Klerman⁵ pointed out that there are individuals who "amplify body sensations", focus on them, misinterpret them and reach the conclusions that they may indicate disease. Costa and McCrae¹² point out that somatizers tend to score highly on measures of neuroticism.

Predisposing factors appear by no means to be solely affected by stressful life events in childhood. In his studies on elderly hypochondriacs, Busse¹³ has noted that contributing factors include recurrent exposure to criticism where there is no possibility of escape, reduction in economic status, loss of spouse and friends, isolation due to socioeconomic factors and deterioration in marital satisfaction.

As Lipowski⁷ notes, sociocultural factors, such as linguistic habits, health beliefs and inhibition of expression, may play a role in somatization.

EPIDEMIOLOGY

While hypochondriasis is seen throughout the life cycle, it is the most frequent somatoform disorder in the elderly. One epidemiologic study¹⁴ shows that 15% of the elderly in the community over the age of 65 reported perceiving that their physical health was poorer than their actual health status. Other investigators¹⁵ have collected data on adults, but not necessarily exclusively in the elderly. Kellner and Sheffield¹⁵ found that 60–80% of physically healthy individuals in the community had at least one physical complaint in 1 week. In their review of the prevalence of "functional complaints" in primary health care, Barsky and Klerman⁵ show that a large proportion of patients (30–80%) who present to the doctor's office do not have evidence of significant physical disease. In their report of the Piedmont Epidemiologic Catchment Area (ECA) study, Swartz *et al.*¹⁶ suggested that somatization symptoms are more prevalent in the rural than in the urban community, that somatization increases with age up to age 65, then tends to drop off, but still remains higher in the elderly over 65 than the age group 18–44. Interestingly, between the ages of 45 and 64, an increase in somatic symptoms is associated with separation, divorce and widowhood. Longitudinal studies of elderly persons living in the community revealed that hypochondriacal episodes are often transient, lasting a few months to several years. The hypochondriacal reaction is often a maladaptive response to social stress¹⁷. Another study¹⁸ concludes that hypochondriasis is less a direct function of stressful life events than of the underlying personality, i.e. a characteristic way of perceiving life events. Patients with transient hypochondriacal reactions are common in a general medical clinic. A recent report¹⁹ notes that, among outpatients confronted with a medical illness, those who are sensitive to somatic sensations and those with personality disorders are more likely to develop hypochondriasis.

TREATMENT CONSIDERATIONS

A number of psychotherapeutic approaches, including individual and group therapy²⁰, have been used in the treatment of hypochondriacal patients. Frequently, individual psychotherapeutic approaches have been based on the treatment methods for depression in those beyond the age of 60 years. This is understandable, because depression is not an unusual feature of hypochondriasis. However, many approaches have certain features in common such as the recognition of the hypochondriac's hostility toward the medical profession and the need to deal effectively with this hostility. Another is avoiding confrontation, specifically the insistence by the therapist that no pathology is present^{17,21}. This is of particular importance to the geriatrician, as hypochondriasis is complicated by the existence of physical disabilities and degenerative disease in the majority of elderly persons.

Hospitalization for extensive medical evaluation is to be avoided¹³ as the experience often increases the hypochondriac's conviction that a "missed" serious illness is likely and adds to the resistance to psychotherapy. Medications may be useful and often do have a transient placebo effect. Drugs should be selected with considerable care to avoid the complications of side effects and addictive qualities.

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Other Neurotic Disorders

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REACTION TO SEVERE STRESS, AND ADJUSTMENT DISORDERS

Before World War II it was generally held that psychiatric patients were constitutionally different from “normals”. During World War II it was observed that previously asymptomatic individuals experiencing unusual environmental stress sometimes suffered from transient psychiatric difficulties. This observation led to a reclassification of psychiatric disorders to allow for behavioral and emotional symptoms in people who would return to their premorbid state with the removal of the unusual environmental precipitant¹. DSM-I and ICD-6 classified these transient difficulties as “gross stress reaction” and “adult situational reaction”; DSM-II and ICD-8 classified them as “transient situational disturbances”. ICD-9 introduced the categories of “acute reaction to stress” and “adjustment disorder”. ICD-10 defines “acute stress reaction”, “post-traumatic stress disorder” and “adjustment disorder”; although DSM-III-R recognized only the latter two of these, DSM-IV recognizes all three.

ACUTE STRESS REACTION

Clinical Features

According to ICD-10, acute stress reaction is a transient disturbance, occurring in persons without apparent mental disorder, in response to exceptional physical and/or mental stress and subsiding in hours or days. The diagnosis should not be made for an exacerbation of symptoms of a diagnosable psychiatric disorder already present, except for accentuation of personality traits. Previous history of another psychiatric disorder does not invalidate this diagnosis. An immediate, clear connection between the stressor and the onset of symptoms should be seen.

Symptoms of this disorder show a mixed and changing picture, with no one symptom predominating for long. They appear within minutes of the stress and resolve rapidly when the stressor is removed or, if the stress remains, symptoms decrease after 24–48 h and are minimal after 3 days.

Typical symptoms include an initial state of “daze”, constriction of consciousness, narrowing of attention, decreased comprehension of stimuli and disorientation. Withdrawal, agitation or overactivity may follow. Autonomic signs of panic (tachycardia, sweating, flushing) are common. Amnesia for the traumatic present may also be present. In the elderly, organic factors and life stage events can be predisposing factors to acute stress reaction². The multiple bereavements which are not uncommon in late life can be the precipitants for acute stress reaction.

DSM-IV differs somewhat from ICD-10 in its diagnostic classification of acute stress disorder. Unlike ICD-10, which requires that symptoms appear within minutes of the stress and diminish to minimal intensity after 3 days, DSM-IV requires symptoms to last a minimum of 2 days and allows for persistence up to 4 weeks. DSM-IV also includes dissociative symptoms not included under ICD-10: a subjective sense of numbing, detachment or absence of emotional responsiveness; derealization and depersonalization. Another DSM-IV requirement is that the traumatic event is persistently re-experienced in at least one of the following ways: recurrent images, thoughts, dreams, illusions, flashback episodes, or a sense of reliving the experience; or distress on exposure to reminders of the traumatic event. DSM-IV also requires marked avoidance of stimuli that arouse recollections of the trauma³.

Differential Diagnosis

The differential diagnosis includes post-traumatic stress disorder (PTSD) and adjustment disorder. PTSD (see below) occurs after a latency period of weeks or longer, while the symptoms of acute stress reaction begin immediately after the traumatic event. The repetitive, intrusive imagery characteristic of PTSD is not usually a feature of the ICD-10 diagnosis of acute stress reaction. DSM-IV, however, does include repetitive intrusive imagery among the features of acute stress disorder. If psychotic symptoms follow an extreme stress, acute (brief) psychotic disorder should be considered. Adjustment disorders are less severe, and longer lasting, than acute stress reactions. Events that precipitate adjustment disorders are also less intense than those responsible for acute stress reactions.

Therapy

By definition, the symptoms of acute stress reaction are time-limited and will resolve without specific therapeutic intervention. Treatment may be requested, however, for intolerable tension or insomnia. For tension, short-term use of benzodiazepines with simple metabolic pathways and short half-lives, such as lorazepam or oxazepam, are safest in the elderly. For insomnia, temazepam or the non-benzodiazepine sedative hypnotic zolpiden⁴ are justified. Families and patients may be reassured that the acute response does not indicate a psychotic decompensation, and that the prognosis for rapid recovery is favorable. Acutely, and in the aftermath of the traumatic event, it is useful to help the patient gain mastery over the trauma⁵ by using a brief treatment model, consisting of fostering abreaction and integration of the event as

quickly as possible, with the expectation that the trauma victim will return to full functioning. Abreaction can be fostered through individual or group psychotherapy⁶.

POST-TRAUMATIC STRESS DISORDER

Post-traumatic stress disorder (PTSD) first appeared in DSM-III but was based on older concepts tied to the history of warfare. Da Costa wrote of "irritable heart" following the American Civil War. In World War I the disorder was known as "shell shock". Early twentieth century psychoanalytic theory called it "traumatic neurosis" and in World War II it was known as "traumatic war neurosis" or "combat neurosis". In DSM-I it was renamed "gross stress reaction", a reaction to great stress in a normal personality. During the peace time between World War II and Vietnam, the category was omitted from DSM-II⁷. ICD-9 defined catastrophic stress and combat fatigue as two diagnoses under the category of acute reaction to stress.

DSM-III defined intrusive re-experience of the trauma, together with emotional numbing, as the central features of PTSD. DSM-III-R placed more emphasis on the avoidance of stimuli associated with the trauma and less on numbing. DSM-IV changed the definition of the trauma to an event where a person experienced, witnessed or was confronted with threatened death or serious injury or threat to physical integrity of self or others. Here, the response to the trauma involves intense fear, helplessness or horror. Also, where DSM-III-R required either numbing or avoidance behavior, DSM-IV requires both³.

ICD-10 criteria more closely resemble those of DSM-III, highlighting the restriction of emotional responsiveness. In ICD-10 the late chronic sequelae of devastating stress, i.e. those manifesting decades after the stressful experience, should be classified under enduring personality change after catastrophic experience².

Clinical Features

The ICD-10 diagnosis of PTSD requires evidence of trauma, or a response to a stressful event or situation of exceptionally threatening or catastrophic nature, likely to cause pervasive distress in anyone. The central symptoms are repetitive and intrusive recollections (flashbacks) or re-enactment of the event in memories, daytime imagery or dreams. The onset follows the trauma with a latency period of a few weeks to months (rarely exceeding 6 months). There may also be a sense of "numbness" and emotional blunting, and avoidance of activities and situations reminiscent of trauma.

Anxiety, depression, suicidal ideation and insomnia are also common in many PTSD patients, particularly with advancing age⁷. PTSD is also associated with alcohol and drug abuse, possibly reflecting attempts to cope with PTSD symptoms. Dissociative symptoms, commonly described in younger PTSD victims, become less prevalent with increasing age^{8,9}.

It remains a subject of debate what factors, if any, predispose individuals to the development of the post-traumatic stress syndrome. Some traumata, particularly the concentration camp experience, are so severe that symptoms are almost universal in survivors. Because retrospective assessment of function before the traumatic event is always colored by the response to the event, correlations are difficult to draw and empirical analyses have been inconclusive¹⁰. Certain personality traits (e.g. compulsive, asthenic) and a previous history of neurotic illness may possibly lower the threshold for manifestation of the disorder².

PTSD can also develop from bereavement. A recent study surveyed surviving spouses 2 months after their spouses' deaths;

10% of those whose spouses died after a chronic illness met criteria for PTSD; 9% of those whose spouses died unexpectedly met PTSD criteria; and 36% of those whose spouses died from "unnatural" causes (suicide or accident) had PTSD¹¹. PTSD can also occur when patients have suffered the "trauma" of having a stroke (9.8%)¹² and upon learning that they have breast cancer (3%)¹³.

Although PTSD symptoms can persist for many years, with increased frequency of symptoms towards the end of life¹⁴, the typical course is one of fluctuating symptoms¹⁵ in many cases. One study, examining current PTSD symptoms in elderly World War II and Korean War prisoners of war (POWs), suggested that severity of exposure to trauma and lack of post-military social support were moderately predictive of PTSD. In this study, 53% of POWs met criteria for lifetime PTSD, with 29% meeting criteria for current PTSD, but for those POWs most severely traumatized, the lifetime PTSD rates were 83%, with current PTSD at 59%¹⁶.

There are two types of PTSD to which the elderly seem susceptible: delayed-onset PTSD and chronic PTSD. In delayed-onset PTSD, patients may exhibit signs of the disorder decades after the trauma, and in chronic PTSD symptoms have been persistent since the time of the trauma.

Delayed-onset PTSD is a reactivation of an old PTSD, with many years relatively free of symptoms, or the first onset of symptoms years after the trauma. In some elderly World War II veterans, media coverage commemorating the 50th anniversary of the end of the war triggered PTSD symptoms¹⁷. Commonly, guilt, distorted memory, emotional numbing, estrangement and feelings of detachment, are seen¹⁸. Patients in this group can present with physical symptoms of cardiovascular, gastrointestinal and musculoskeletal diseases¹⁰. Generally, the onset of severe symptoms can be linked to a profound recent life event, such as death of a wife, job retirement or loss of physical integrity from illness¹⁸. Most often, the contemporary precipitant reawakens emotions and perceptions from the original trauma. Holocaust survivors and prisoners of war have been noted to begin displaying symptoms of PTSD after admission to nursing homes, where they re-experience a loss of freedom and autonomy. World War II veterans found the loss of physical integrity due to somatic illness particularly upsetting, since it evoked memories of a traumatic period when their physical integrity was in jeopardy⁹.

Differential Diagnosis

Although adjustment disorders also occur in response to life events, these events are in the normal range of human experience, unlike the extraordinary traumata responsible for PTSD. Specific features of numbing and flashbacks do not occur, and adjustment disorders, by definition, do not last more than 6 months. Acute stress reaction is characterized by a more variable clinical picture that resolves within days.

While anxiety and depression are common features of PTSD, generalized anxiety disorder and phobic disorder have anxiety as a more specific and central symptom. Major depression is marked by deep and persistent mood disturbance, usually with loss of reactivity; dysthymia results in chronic, indolent dysphoria. None of these disorders includes the specific symptom of intrusive recollections.

Therapy

The signs and symptoms of post-traumatic stress disorder include distorted expectations and perceptions, mood disturbances, psychophysiological symptoms and social withdrawal. Thus,

common sense dictates, and empirical data confirm, that multimodal treatment is most advisable. Psychosocial intervention and pharmacotherapy each has its place. At all ages, the psychotherapy of PTSD starts with the retelling of the story of the events before, during and after the traumatic episode. The goal is to integrate the experience with the person's life history; the method is to frame the events from the perspective of the intact self, rather than leave them relegated to the weakened self of the past. It is particularly important in older patients, given the chronological distance from the event, to differentiate objective properties of the trauma from the fantasy attributions it accumulates over time¹⁹.

Group therapy has been found to be particularly useful. Matching patients by age and by setting of trauma enhances feelings of understanding and group identification²⁰. The focus of the group is on recurring memories, and benefits patients by relieving long-held guilt through objective evaluation of the traumatic incident, as well as enhancing their ability to tolerate life stressors^{18,21,22}.

Antidepressants can offer symptomatic relief by diminishing dysphoria, intrusive thoughts, insomnia and nightmares. In particular, selective serotonin reuptake inhibitors (SSRIs) can be effective, especially in reducing avoidant symptoms^{24,25}. β -Adrenergic blocking agents have been used in younger PTSD patients for relief of symptoms of autonomic arousal, tremors and startle reactions²⁶. Older patients, however, are less likely to display a clinical profile of hyperarousal, and are more susceptible to the cardiovascular complications and organic mood disorders associated with adrenergic blockade. Benzodiazepines should be avoided as much as possible, since they can cause paradoxical excitation and frequently induce subtle cognitive impairment in aging individuals²⁷.

ADJUSTMENT DISORDER

The diagnosis of adjustment disorder refers to a state of subjective distress or emotional disturbance, interfering with social functioning or performance, arising in a period of adaptation to a significant life change or subsequent to a stressful life event. It is assumed that the condition would not have arisen without the stressor. In ICD-10, onset is usually within 1 month of the stressor; in DSM-IV, it can be within 3 months of the stressor. In both ICD-10 and DSM-IV, duration of symptoms does not exceed 6 months, except in the case where the stressor is chronic (e.g. a chronic general medical condition) or the stressor has enduring consequences (e.g. the financial and emotional difficulties resulting from a divorce)³.

Clinical Features

Symptoms of adjustment disorder may include: depressed mood, anxiety, worry, impairment in performance of daily routines and inability to cope or plan ahead. Adjustment disorders can be specified as brief depressive reaction, prolonged depressive reaction, adjustment disorder with predominant disturbance of other emotions, adjustment disorder with predominant disturbance of conduct, or adjustment disorder with mixed disturbance of emotions and conduct.

The precipitating events for adjustment disorders can affect social network or values, and may involve the individual, his group or community. Common events causing such symptoms in older patients include physical illness or injury, placement in a nursing home and retirement. The events, while subjectively profoundly meaningful, are of considerably smaller magnitude than those precipitating acute stress reaction and PTSD. Individual predisposition and vulnerability to these stressful life

events thus plays a greater role in the occurrence of adjustment disorders. Poor pre-stressor social and coexisting physical problems²⁸, current dementia²⁹ and a history of a past psychiatric disorder³⁰ all increase vulnerability to adjustment disorders.

Therapy

The cornerstone of treatment for adjustment disorders is focal psychotherapy. Based on a psychodynamic understanding of emotions and behavior, focal therapy identifies the most specific nidus of current distress and views it in the context of the patient's core conflicts or deficits. The therapy is of relatively brief duration, usually 6–20 sessions. The major techniques employed are clarification and confrontation³¹.

Quite frequently, the precipitating event can be framed as a narcissistic threat or injury. In psychotherapy, the patient will come to view the therapist as a self-object, looking for restoration of the self-esteem provided by the lost function, role or friend. The therapist helps restore the wholeness of self by allowing the patient to modify his/her expectations of him/herself and environment³².

DISSOCIATIVE AND CONVERSION DISORDERS

In the last three decades of the nineteenth century, dissociation was studied extensively by Janet and conversion by Freud. DSM-I incorporated the concepts of dissociation and conversion into its classification scheme. Conversion reaction was assigned to hysterical neurosis, and amnesia was placed in the category of dissociative reaction. In DSM-II they were united under the heading of hysterical neurosis, but divided into conversion type and dissociative type. In DSM-III, DSM-III-R and DSM-IV the two conditions were renamed and separated once again. Hysteria, conversion type, became conversion disorder and was assigned to somatoform disorders. Hysteria, dissociative type, was expanded into the dissociative disorders³³. ICD-10, however, continues to contain both under the heading of dissociative disorders.

DISSOCIATIVE AMNESIA

Dissociative amnesia is characterized by loss of memory, usually of important recent events, that is too great to be explained by ordinary forgetfulness or fatigue; and amnesia, either partial or complete, for recent events that are of a traumatic or stressful nature. The amnesia is usually partial and selective. The extent and completeness of the amnesia varies from day to day and between inquirers, but a persistent common core cannot be recalled in the waking state. Complete, generalized amnesia is rare and is usually part of a dissociative fugue. Affective states in amnesia are varied but severe depression is rare. Perplexity, distress and varying degrees of attention-seeking behavior may be evident, but calm acceptance is also sometimes striking. Purposeless local wandering may occur, but is rarely accompanied by self-neglect and rarely lasts more than a day or two. Often in dissociative amnesia, new learning is preserved³⁴. Disturbing external circumstances causing despair or anxiety may predispose an individual, but a single event is usually at the center of the syndrome.

Dissociative amnesia is uncommonly reported in the elderly, but has been seen in World War I combat soldiers³⁵ and soldiers in other conflicts. In most patients the amnesia is short-lived, 75% of cases lasting between 24 h and 5 days³⁶. Its features resemble those of more frequently observed disorders. Organic amnesia is usually anterograde³⁴. In postconcussional syndromes there may

be a combination of hysterical and organic amnesia that can be difficult to untangle. In dementia, memory loss is seen in the context of global cognitive impairment, which is stable over a period of weeks to months. The syndrome of pseudodementia also features variable memory impairment, but affective disturbance, usually severe depression, is evident³⁷.

DISSOCIATIVE FUGUE

Fugue exhibits all the features of dissociative amnesia, plus an apparently purposeful journey away from home or place of work during which self-care is maintained. A new identity is assumed and organized travel may be undertaken to places previously known and of possible emotional significance. Although there is retrograde amnesia during the fugue, behavior during fugue is normal.

A severe precipitating stress is almost universal as a precipitant of dissociative fugue. Times of marital discord, financial difficulty, major role change or personal loss may precede the fugue. Depressed mood is frequently present before fugue symptoms are displayed³⁴.

Fugue is rare in elderly people. Because its features, with the exception of travel, are identical to those of dissociative amnesia, it has been proposed that the two disorders be considered as one.

Treatment

Therapy for dissociative amnesia and dissociative fugue is virtually identical. Patients usually seek treatment after the amnesic period has ended. They desire help in recovering memory of events during the fugue. Hypnosis and short-acting barbiturates have been used to reconstitute repressed memories, although typically they return spontaneously. Psychodynamic psychotherapy has been used to facilitate resolution of conflicts that lead to fugue states. This treatment may decrease the vulnerability of the patient to dissociate in future times of stress³⁸.

DISSOCIATIVE DISORDERS OF MOVEMENT AND SENSATION (CONVERSION DISORDERS)

In conversion disorder, there is a loss or alteration in movements or sensations (usually cutaneous) in a patient presenting as having a physical disorder. No somatic condition can be found, however, that explains the symptoms. Instead, the symptoms represent the patient's concept of the physical disorder, which may be at variance with physiological or anatomical principles. Here, mental state and social situation suggest that disability resulting from the loss of function is helping the patient to escape an unpleasant conflict, or helps the patient to express dependency or resentment indirectly. Conflicts may be evident to others, but the patient often denies their presence and attributes distress to the physical symptoms or the resulting disability.

In making the diagnosis it is essential that: (a) evidence of a physical disorder is absent; and (b) sufficient knowledge of the psychological and social setting and personal relationships of the patient allows a convincing formulation of the reasons for the disorder.

Predisposing factors to conversion disorder are premorbid abnormalities of personal relationships and personality. Also, close relatives or friends may have suffered from physical illness with symptoms resembling the patient's. A few patients establish a repetitive pattern of reaction to stress by production of these disorders, which can continue into middle and old age².

The most important differential diagnosis is the group of somatoform disorders (although DSM-IV classifies conversion disorder as a somatoform disorder instead of a dissociative disorder). In the latter, the patient's presentation centers around persistent requests for medical attention and pervasive concern with the perceived medical disorder; patients with conversion disorder are much more likely to take their presumed illnesses in stride. Conversion disorders generally begin in adolescence and young adulthood, and occur in single or recurrent episodes with substantial remission. Somatoform disorders may not increase in prevalence with increasing age, but they tend to assume the quality of a pervasive character style with little remission.

Most conversion disorders remit with non-specific, supportive interventions. Hypnosis, anxiolytics and behavioral relaxation exercises may be helpful. Also, psychotherapy aimed at helping the patient recognize and cope with the psychosocial stress that provoked the symptom can be impressively beneficial if the patient can be engaged in a cooperative alliance of therapeutic curiosity.

The prognosis of conversion disorder is generally good, since conversion symptoms are of short duration with abrupt onset and resolution. A few become chronic, and some recur, most commonly when the precipitating stress is chronic or recurrent, when there is other psychopathology or when there is marked secondary gain.

NEURASTHENIA (FATIGUE SYNDROME)

Historical Perspective

George Beard introduced the term 'neurasthenia' in 1869. He viewed neurasthenia as a physical illness due to loss of nerve strength. Janet differentiated psychasthenia from neurasthenia. Freud similarly separated anxiety neurosis, a "psychoneurosis", from neurasthenia, an "actual neurosis" he attributed to misdirected libidinal energy.

In World War I, the syndrome was defined by the term "shell-shock"; in World War II, "operational fatigue". Although it remains in ICD-10, the diagnosis of neurasthenia was deleted from DSM-III and replaced by dysthymia. In the USA the symptom cluster known as chronic fatigue syndrome is almost identical to the current ICD classification of neurasthenia³⁹⁻⁴¹.

Clinical Features

Neurasthenia is characterized by persistent, distressing complaints of fatigue after mental effort, or complaints of bodily weakness and exhaustion after minimal physical effort, along with at least two of the following: muscular aches and pains, dizziness, tension headaches, sleep disturbance, inability to relax, irritability or dyspepsia. If autonomic or depressive symptoms are present, they cannot be sufficiently persistent and severe to fulfill the criteria for any more specific disorder².

Differential Diagnosis

Differential diagnosis includes primarily major depression and somatoform disorders. In the elderly it is especially important to rule out depression, since somatic complaints and fatigue are common presentations of depressive disorders in late life. Physical symptoms with no demonstrable organic pathology are the essential features of somatoform disorders. However, these complaints do not include the specific physical symptoms of fatigue or exhaustion found in neurasthenia.

Therapy

Specific treatment for neurasthenia has not been established. Given the high likelihood, particularly in old age, that the neurasthenic picture is a manifestation of a mood disorder, treatment with antidepressant medication and psychotherapy, as for depressive conditions, is generally warranted.

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Part I

Personality Disorders

Personality Disorders: Aetiology and Genetics

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According to DSM-IV¹, “A Personality Disorder is an enduring pattern of inner experience and behavior that deviates markedly from the expectations of the individual’s culture, is pervasive and inflexible, has an onset in adolescence or early adulthood, is stable over time, and leads to distress or impairment” (p. 629). Since the first edition of this chapter was published, there has been an ever-expanding body of knowledge about personality disorder (PD) in older adults. Notably, there has even been the publication of the first book² solely devoted to PD in older adults. However, as we shall see, there remain many unanswered questions spawned by thorny conceptual and methodological quandaries in this controversial area.

PD in older adults is an important area of study for a number of reasons. First, since PD affects the way an older adult copes with life, individuals with specific PDs may be less able to successfully negotiate age-related losses (e.g. obsessive-compulsive, dependent) or the interpersonal compromises necessary for peaceful institutional living (e.g. borderline, narcissistic). Second, PD can influence the presentation of Axis I symptomatology, frequently generating complicated assessment dilemmas. For example, disruptive behavior in the nursing home may camouflage the fact that the person is suffering from a depression that is exacerbating premorbid antisocial personality features. Third, just as for young adults, the presence of PD should modify treatment strategies and prognosis of co-morbid Axis I disorders in certain geriatric settings.

This chapter will summarize what is known about the etiology, genetics, epidemiology, assessment, prognosis and research implications of PD in older adults.

AETIOLOGY AND GENETICS

Reviewing PD necessitates some understanding of the complexity of the terms “personality disorder” and “personality”, as well as their classification. Personality can be conceived as two interactive elements, representing “nature” and “nurture”: temperament, a reflection of a genetically determined, constitutional disposition; and character, made up of learned attributes, which begin coalescing in early childhood, reflecting culture, norms and upbringing. Studies have focused on each of these elements in order to clarify the etiology of personality and PD, but these investigative efforts have been rendered more complicated by different measurement approaches.

Concerning PD among the elderly, it is helpful to focus on dimensional aspects of personality as well as the categorical diagnosis of PD. Because of the numerous biological changes that occur with aging, a major question for the geriatric clinician concerns the stability of personality traits throughout the aging process.

Most researchers agree that there is no uniform, stereotypic change in personality traits in late life. Cross-sectional and longitudinal investigations of personality across the life cycle³ report the general stability of individuals’ major traits. However, it is also clear that certain adults do change for the better or worse⁴, perhaps due to some of the unique challenges they face over the lifespan.

Late-life theorists have made important contributions to conceptualizing those challenges that promote personality development. Erikson’s⁵ crisis of “integrity vs. despair” marks the culmination of his psychosocial framework. The challenge of this final stage is the individual’s acceptance of the integrity of his/her self; failure results in a fear of death and an inability to find meaning in the life cycle. Erikson felt that the successful resolution of this stage depends on an adequate resolution of previous crises. From the point of view of personality, failure here may also trigger global dissatisfaction and lowered self-esteem. Erikson’s developmental stages, intended for a practitioner audience, were not rigorously defined and were based on clinical and theoretical considerations rather than research data.

Levinson *et al.*’s⁶ developmental scheme is based on their research on the life cycle. They view development as a series of eras and cross-era transitions. Older adults are seen from the point of view of the late adult transition and late adulthood. The late adult transition (age 60–65) is marked by major changes in role structure, physiology and intrapsychic challenges. It is possible that adaptation to these challenges provokes changes in personality structure (i.e. a decrease in authoritarianism; locus of satisfaction shifting from the self to others). Haan and Day⁷ traced the relative prominence of different personal dimensions (such as information processing, inter-personal reaction, manner of self-presentation and responses to socialization) throughout the life cycle from adolescence onwards. Although most dimensions were stable, some changed in an orderly manner with age. Still others changed in a stage-wise manner. The authors believed that their results fitted an Eriksonian developmental model with an essential “sameness” of major personality functions, punctuated by times of regress before progress and longer periods of orderly change.

The investigation of constitutional factors that contribute to PD is affected by a number of methodological difficulties. Few studies have looked specifically at DSM Axis II criteria; many have looked at other variables, such as neuroticism and sociability, which cannot be compared well with PD as defined by DSM.

The most intensive studies of familial factors have been conducted on those with antisocial PD. Crowe⁸ summarized data from twin studies in strong support of a heritable component

for antisocial PD. The mean concordance rate for monozygotic twins for the disorder was double that of dizygotic twins (68% vs. 33%). Adoption studies^{9,10} have also found “temperamental” factors to be a major determinant for antisocial PD. Family studies find increased rates of substance abuse, antisocial PD and other psychopathology in first-degree relatives of felons, many with the diagnosis of antisocial PD¹¹.

Schizotypal PD has been well examined from a genetic perspective and owes much of its appearance as a diagnosis to that association. Schizotypal PD has a significant heritable relationship to schizophrenia. Kendler *et al.*¹² were unable to find an equivalent environmental association. Torgersen¹³ confirmed the genetic link between schizoid and paranoid features of schizophrenia and schizotypal PD.

The final disorder that has yielded studies on heritability is borderline PD. A credible familial association has been found between borderline PD, affective disorders and other PD¹⁴. Previous associations between “borderline states” and the schizophrenic spectrum have been difficult to interpret because of differing definitions of “borderline states” and the relationships of these definitions to current DSM-IV criteria for borderline PD. Individuals with a history of impulsive or affective instability, which more closely fit present borderline PD criteria, may not have a greater prevalence of schizophrenia in their relatives¹⁴.

In summary, there is clear evidence of heritability for some PD but much that remains unexamined. As Clarkin, Speilman and Klausner¹⁵ note in their conceptual overview of PD in the elderly, the research on the relationship of genetic factors to PD is sparse, and speculations concerning the neurobiology of PD remains premature. The remaining “nurture” factors also comprise a significant component, but have yet to be clarified.

Neurobiological studies have been very rare. EEG slow-wave activity has been associated with antisocial PD. Borderline patients have a significantly higher percentage of marginal EEG abnormalities than controls¹⁶. They also show changes in cerebrospinal 5-hydroxyindole-acetic acid (5-HIAA), a measure of serotonin activity that is correlated negatively with measures of aggressivity. Finally, schizotypal and borderline PD have been linked with low platelet monoamine oxidase (MAO) activity^{17,18}. Individuals with schizotypal PD may also show impaired smooth pursuit eye movement.^{19,20}

ASSESSMENT

Assessment of PD requires an examination of variables that may be difficult to measure, especially in older adults. There are, for instance, problems in obtaining a reliable diagnosis. Since all the standardized instruments used to measure PD have been developed for a younger population, these devices thereby require more administration time for older adults (since they have longer personal histories). Consequently, there is no “gold standard” of diagnosis for PD in older adults. Molinari *et al.*²¹ studied geropsychiatric inpatients with depression, and found general discordance between patient self-report, family informant ratings, social worker evaluations, and consensus case conference categorical diagnosis of PD. It appears that there are varied perceptions of an individual’s personality, all of which should be taken into account for a comprehensive evaluation of Axis II pathology.

From a clinical perspective, PD is quite common in medical practice, yet infrequently recognized. Mental health professionals are loathe to diagnose PD, particularly in old age, due to concerns over pejorative bias, pessimistic belief in Axis II changes, managed care reimbursement biases, and focus on medical or Axis I pathology (particularly cognitive impairment) in old age. Often the PD patient presents with unrealistic expectations,

complaints and demands, and an inappropriate interpersonal stance. Essential features of PD, regardless of age, will include: maladaptive behavior as a lifestyle; inflexibility in managing interpersonal situations; multiple physical, social and interpersonal problems; and externalization of these problems in the absence of psychosis²². Yet these same features are sometimes accepted, erroneously, as part of the aging process. In a geriatric setting, somatic presentations of PD are common, which can complicate “teasing out” true co-morbid medical/cognitive problems from personality dysfunction. Patients with PD frequently present with chief complaints of “bad nerves”, sleeplessness, non-specific requests for help and pain syndromes. In addition, Axis I disorders can confound appropriate diagnosis and confuse the long-term picture. For example, evidence suggests that older depressives are significantly more likely to have lifetime personality dysfunction than controls^{23,24}. Therefore, it behooves the clinician to ascertain whether a pattern of historically maladaptive personality traits exist in the older patient with depression. Treatment of the depression may very well result in a symptom picture reflecting a baseline PD, rather than a depression in partial remission.

EPIDEMIOLOGY

Some early anecdotal reports suggested that personality characteristics become uniformly less harsh with age^{25,26}. Other clinicians working with older adults believed that the “high-energy” PDs (e.g. Cluster B) mellow, while the “low-energy” PDs (e.g. Cluster C) may be aggravated by the aging process²⁷⁻²⁹. DSM-IV¹ states that, “Some PDs tend to become less obvious or remit with age, whereas this appears to be less true for some other types” (p. 632). This statement underscores some of the more recent empirical work in this area, and contrasts with the more benign general appraisals of DSM-III³⁰ and DSM-III-R³¹, which merely note that PD becomes “less obvious” with age. Early research yielded wide variability in PD prevalence rates due to inadequate definitions of PD, non-standardized measures, and different samples of older adults. With the introduction of DSM-III and the development of instruments tied to DSM-III criteria, some consistent findings have emerged. This section on epidemiology will therefore largely focus on studies using standardized measures, and will be divided into community, institutional, outpatient and depression studies.

Community Settings

In community settings, two studies^{26,32} compared young and older adults utilizing the Coolidge Axis II Inventory. Coolidge *et al.*²⁶, found a greater need for organization and more restricted affect in older adults, while Segal *et al.*³² found that older adults were significantly higher on obsessive-compulsive and schizoid PD, but lower on the antisocial, borderline, histrionic, narcissistic and paranoid scales. Ames and Molinari³³ used the Structured Interview for Disorders of Personality scale (SIDP-R) and detected a trend of less PD in older adults, with significantly fewer older adults meeting the criteria for more than one PD. Cohen *et al.*³⁴ used the Structured Psychiatric Examination and found that those over the age of 55 were less likely (6.6% vs. 10.5%) to have PD, due to a three-fold decrease of Cluster B PD in older adults. These data documenting personality “mellowing” in older adult community samples are in stark contrast to the results of a more recent study by Segal *et al.*³⁵, who found that a high number (63%) of very old ($X=76.2$ years) community-dwelling adults met PD criteria by self-report. However, this study was potentially flawed by failure to take into account the cognitive

and sensory status of this aged sample, many of whom may have misunderstood the questions or exhibited executive deficit-related personality alterations.

Institutional Settings

Early PD prevalence rates in nursing home settings were reported to be 12–15%^{36,37}, while for geropsychiatric inpatients, PD estimates were more variable (7–55%). In a large sample of hospitalized male veterans, Molinari *et al.*³⁸ conducted a cross-sectional investigation of personality changes across different age groups for those clinically diagnosed with PD. Older adults with PD were more responsible and less impulsive, paranoid, energetic and antisocial than young adults diagnosed with PD. Kunik *et al.*³⁹ studied 547 older psychiatric inpatients, and found that a consensus case conference diagnosis of PD varied widely, depending upon the specific co-morbid Axis I diagnosis (e.g. 6% for patients with an organic mental disorder, but 24% for those with depression). Only two studies of geropsychiatric institutionalized patients utilized standardized instruments. Molinari *et al.*⁴⁰ used the SIDP-R and found that older adults had PD rates similar to those of a young adult comparison sample; however, older adults were less likely to meet criteria for more than one PD, and clinical diagnoses yielded fewer PDs than the SIDP-R. Likewise, Coolidge *et al.*⁴¹ used the Coolidge Axis II Inventory and found similarly high PD rates among young (66%) and old (58%) chronically mentally ill patients, but the younger group was more likely to be specifically diagnosed with antisocial, borderline, and schizotypal PD.

Outpatient Settings

The findings from the lone study conducted with a structured PD scale in a geropsychiatric outpatient setting are consistent with the latter inpatient studies. Molinari and Marmion⁴² found that older adults were less likely to meet the criteria for more than one PD than younger adults, and clinical diagnosis again yielded fewer PDs than the SIDP-R.

Depression

One area of intense study has been the relationship between PD and depression in older adults. Kunik *et al.*²⁴ studied 154 depressed older inpatients and identified 24% with co-morbid PD, while Molinari and Marmion⁴³ determined that 63% of depressed geropsychiatric outpatients met PD criteria. Thompson *et al.*⁴⁴ found that 33% of depressed older adults who were being treated with psychotherapy in a geropsychiatric outpatient clinic met DSM-III PD criteria. In a study investigating the relationship between PD and functioning in acutely depressed older psychiatric patients, Axis II pathology was found to be associated with greater disability and more impaired social and interpersonal functioning⁴⁵. In their review of the literature on PD in older adults, Agronin and Maletta⁴⁶ posit that PD in late life may be intrinsically related to Axis I pathology, particularly major depressive disorder.

Summary of Epidemiological Studies

In an attempt to lend clarity to the burgeoning literature on PD in older adults, Abrams and Horowitz⁴⁷ conducted a meta-analysis of the most methodologically sophisticated epidemiological studies. They inferred a PD prevalence rate of 10% (with a

range of 6–33%) for those over the age of 50, and concluded that, “at this time the literature neither supports nor contradicts previous suggestions of an age effect”. However, these authors remark that the bulk of the evidence supports, at least for certain PDs, a decline in frequency and intensity with age. The cause for this decline is one of the most controversial and debated topics in the literature on PD in older adults. Four main reasons have been postulated.

First, there is a general mellowing of the “high-energy” Cluster B PDs due to biological (reduced testosterone in males) and developmental changes (those with PD finally master a single interpersonal strategy to manage stresses). This accounts for the consistent result that older adults are less likely to meet the criteria for more than one PD, and is also supported by the recent study of Segal *et al.*³², who discovered lower levels of dysfunctional coping styles in older adults.

Second, the decline in “high-energy” PD relates to the greater mortality rates of those with Cluster B PD in their younger years. Older adults with PD are thereby a selective sample of less extreme PD “survivors”. Third, PD is generally underdiagnosed, particularly in older adults, where cognitive and medical causes are emphasized or personality disturbance (avoidance, dependency, lability) is viewed as normal⁴⁸.

Fourth, the decline in PD with age is a methodological artifact, since some DSM criteria are age-insensitive. For example, occupational and vocational impairment are often irrelevant to older adults. From this point of view, there really is no true decline in PD rates with age, just a change in form that is inadequately assessed. These so-called “geriatric variants”⁴⁹ reflect the more subclinical, non-specific or age-relevant PD traits that account for PD NOS (not otherwise specified) to be diagnosed with particular high frequency in older adults. The construction of a new geriatric nosology has been proposed to accommodate the late life changes in Axis II pathology^{46,47,49}. Such re-classification will need to: (a) reconsider the diagnostic requirement that maladaptive PD behavior be rooted so early in young adulthood; (b) routinely address Axis II pathology in the context of more acute Axis I symptomatology; and (c) integrate age-related developmental, medical (Axis III) and psychosocial/environmental stressors (Axis IV) with Axis II manifestations⁴⁶.

PROGNOSIS

Unfortunately, only a few seemingly contradictory studies have investigated the prognosis of PD in late life. In two separate studies of geropsychiatric outpatients, PD was found to be a poor prognostic sign for the psychotherapeutic treatment of depression^{44,50}. However, Molinari⁵¹ examined the 1 year relapse rates for 100 male geropsychiatric inpatients and found no significant differences for those diagnosed with and without PD. Consistent with Kunik *et al.*'s²⁴ finding that PD diagnosis had no impact on the acute response of inpatient treatment for depression with older adults, no differences were found in relapse rates for a subgroup of depressed inpatients with and without PD⁵¹. It appears that in inpatient geropsychiatric settings, Axis I symptomatology overrides Axis II pathology as an outcome predictor, probably related to the complex combination of medical, cognitive and psychiatric symptoms often observed in those older patients needing acute care.

RESEARCH IMPLICATIONS

Base rates mandating prohibitively large sample sizes render it difficult to make valid statements concerning age-related changes for most individual PDs. To expand our current inadequate

knowledge base, research should concentrate on PD clusters and PD with older adults must be conducted using reliable scales tied to DSM-IV criteria across varied settings (e.g. medical, nursing home, psychiatric inpatient, psychiatric outpatient) populations (e.g. old-old, females, multicultural) and methodologies (e.g. longitudinal, cross-sectional). The inclusion of age-graded criteria for Axis II and perhaps also Axis I should be investigated. At the very least, since all the PD scales currently in use have been developed for younger adults and have yielded generally modest reliability ratings for most PD categories, validity studies must be conducted specifically with older adults⁴⁸. Such studies must take into account the effect of cognitive status and medical problems on the responses of older adults. In this way, research concerning the psychological/developmental history of individuals with PD will generate valuable prognostic and treatment data. Research should also explore those PD features underlying late onset Axis I disorders. Hopefully, more sophisticated methodologies will probe the relative and combined merits of pharmacotherapy and different types/modes of psychotherapy (e.g. cognitive-behavioral vs. interpersonal; group vs. individual). Heuristic models of the interrelationship between the underlying neuropsychiatric and biological substrate of temperament, genetic factors and psychosocial changes in personality with aging must be generated. Finally, the ethical and clinical issues involved in the management of "difficult personality-disordered" patients in hospital, rehabilitation and nursing home sites should be explored, so that the needs of individuals can be balanced with those of families, staff and patient/resident peers⁵²⁻⁵⁴.

SUMMARY

Over the last decade, there has been an increasing amount of research on PD in older adults using structured scales tied to DSM criteria. Major findings are:

1. Although still common in older adults, Cluster B pathology is less prevalent than in the young adult population; Cluster C pathology may be relatively more prominent in older adults.
2. Older adults are less likely to receive more than one PD diagnosis, suggesting that they may finally develop one main coping strategy to fulfill their interpersonal needs.
3. There may be an age-related mellowing of the "high-energy" personality characteristic of individuals with PD, and/or there are "geriatric variants" of PD not tapped by DSM.
4. There is a positive association between depression and PD diagnosis.
5. PD in older adults is prognostically useful in outpatient settings, where the Axis I symptomatology is less severe.
6. There are poor concordance rates of PD diagnosis between clinical examination, structured interviews and self-reports, suggesting the need for data collection from a variety of sources.

Although age-related changes in PD expression are probably in the less volatile and impulsive direction, novel PD manifestations can still create a significant burden in stressful caregiving contexts for family members, friends, healthcare professionals and administrators of institutions attempting to support a flawed but vulnerable older adult. Future research guided by conceptual advances promises to yield exciting progress in assessment and treatment.

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Theoretical and Management Issues

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The elaboration by DSM-III¹ of a domain of pathological personality typologies into the well-known Axis II personality disorders has stimulated an outpouring of clinical research. While most of this research has reaffirmed the usefulness of the personality disorder concept, substantial theoretical and methodological problems persist. For example, there is considerable heterogeneity of symptoms within individual disorders and overlap among different disorders². It can be argued that the Axis II personality disorders to some degree lack both convergent and discriminant properties. Moreover, the theoretical underpinning of personality disorders is far from clear. Are these entities derived from interactions of heritable traits, as suggested by Cloninger³, or are they based on a pathological intrapsychic organization, as suggested by Kernberg⁴? Alternatively, are personality disorders better understood in interpersonal or learning-behavior contexts? It has proved possible to apply each of these theoretical models; personality disorders are, by definition, relatively enduring patterns of dysfunctional behavior that permeate the entire person, are not due to a single abnormality and lead to pain and suffering of people in the patient's world, sometimes more than in the patient him/herself. It is thus not surprising that a variety of explanatory models have been used in efforts to understand these disorders.

Very little attention, however, has been directed to the existence of personality disorders in the elderly and their clinical management, despite the considerable interest in long-term follow-up studies of borderline personality disorder patients^{5,6}. In part, this neglect may reflect the original perspective of DSM-III and DSM-III-R⁷, that personality disorders are largely attenuated by middle- and old age. Probably another reason is the ongoing uncertainty about how best to assess personality in the elderly. Aging imposes complexities on the already difficult study of personality, adding the confounding factors of brain changes, environment and time. Finally, concepts of normal and pathological adult personality functioning may not be relevant to old age.

In this chapter the topic of personality disorders in old age will be approached first with a discussion of age-related personality change, followed by a review of methodological and clinical issues, including comments on the assessment of DSM personality disorder criteria in the elderly. It will conclude with a discussion of treatment and management.

THE AGING PERSONALITY: DO PEOPLE ACTUALLY CHANGE?

Although the aging personality has been the subject of extensive speculation^{8–10}, there has been no closure on the question of

whether, or in what ways, people change in their dealings with the world; it is still not known whether change or persistence predominates over time. The literature in this area to date can be divided along theoretical and methodological lines into dimensional, categorical and psychoanalytic–developmental models, each with a different set of assumptions.

Using mostly dimensional approaches, conceived as continua of stable traits along which all individuals lie, the psychological literature suggests a picture of progressive changes in motivations and values over the adult years. For example, there appear to be steady increases from early to middle adulthood in self-confidence, independence, humanitarian concerns and personal rigidity^{8–9}. Large-scale investigations of medical patients and normal subjects have been carried out using the Minnesota Multiphasic Personality Inventory (MMPI)^{11,12}; older adult medical patients tended to score higher on scales measuring introversion, concern with health, immaturity and depression than younger adult medical patients. A decline in criminality with advancing age has also been well documented¹³. Together, these data suggest that traits related to a quiet, inner-directed attitude most typify the aging personality. The MMPI data in particular seem to lend support to the concept of disengagement, or gradual withdrawal from productive activity, as the pre-eminent social-psychological model for age-related personality changes¹⁴. The reduction in sociopathy and attenuation of hostile aggressive traits have also been highly consistent MMPI findings^{1,12}.

However, such dimensionally-modeled studies have not pointed conclusively to the existence or direction of age-related personality changes. For example, the MMPI data referred to above are based on large-scale cross-sectional studies involving thousands of medical patients and normal adult subjects, but have not been confirmed in longitudinal studies; the longitudinal MMPI studies have instead emphasized the stability of personality profiles of the individuals over time^{15,16}. Similarly, dimensional scores on the Eysenck Personality Inventory Psychoticism (P), Extraversion (E) and Neuroticism (N) subscales show substantial persistence over 30 year periods⁷.

Categorical models, such as the DSM-IV Axis II personality disorders, are operationally defined typologies of personality psychopathology, or clinical phenotypes. These models have been relatively little used to examine the aging personality. At the present time, for example, there is scarce cross-sectional information on DSM-IV personality disorders in the older age groups and virtually no longitudinal studies of personality disorder patients that track outcome as far as senescence. However, based upon cross-sectional regression analysis of the relationships of different personality disorder traits with age, Tyrer¹⁸ has suggested that personality disorders can be divided into mature and immature

categories. The mature disorders include obsessive-compulsive, schizotypal, schizoid and paranoid; these personality disorders show more stability and less variation with age than others. Schizotypal personality disorder, for example, would be expected to behave more like schizophrenia from the longitudinal perspective because of its spectrum and genetic relationships to the Axis I disorder⁵⁻⁸. The immature or flamboyant personality disorders include the borderline, antisocial, narcissistic, histrionic and passive-aggressive categories; these personality disorders may be more evident in younger individuals and may have earlier onset than mature personality disorders. In this scheme, mature personality disorders consist primarily of the Cluster A (odd-eccentric) disorders plus obsessive-compulsive disorder from Cluster C (anxious-fearful); immature personality disorders consist primarily of Cluster B (dramatic-emotional) disorders, plus passive-aggressive disorder from Cluster C.

Other authors, notably McGlashan⁵ and Stone⁶, have commented, on the basis of follow-up data, that the florid borderline symptomatology seen at index admissions substantially declines by the second decade of follow-up. Both male and female patients appear to advance occupationally and globally, as well as symptomatically, with time⁵. Tyrer¹⁸ also reviewed data suggesting that patients with personality disorders have higher mortality by suicide than other psychiatric patients for a period of 5 years from diagnosis, after which differences in suicide rates between personality disorders and other psychiatric patients become negligible. These mortality data have been interpreted by Tyrer to support a maturation hypothesis, especially for antisocial and other immature personality disorders, in which impulsiveness and suicide become less likely over time. There are also suggestions that mature personality disorders are more frequent in the geriatric population than immature disorders^{19,20}.

Thus, it is possible to speculate about a "flattening" over time of at least some specific types of personality disorder symptomatology, which might explain why geriatric patients seem to have diffuse Axis II symptomatology, with relatively fewer full diagnoses of personality disorder^{19,20}. In the case of borderline personality disorder, the assumption that symptoms are enduring and inflexible has been challenged; this disorder might now be viewed as a state of delayed maturation that improves with time, rather than as a set of chronic defects. However, even in borderline personality disorder, the improvement may not be uniform across different spheres of functioning. McGlashan⁵ has suggested that borderline patients may improve considerably in occupational or instrumental functioning, yet never develop satisfactory personal relationships.

Moreover, aging does not necessarily imply linear reduction in severity of the immature personality disorders. An epidemiological study of personality disorders conducted by Reich *et al.*²¹ found the dramatic cluster to be described by a reverse J-shaped curve, in which core traits decline to age 60, then take a slight upturn. Geriatric clinical experience also suggests that there are some individuals who have relatively mild personality dysfunction in young and middle adulthood, but in old age develop a marked and persistent worsening of these trends²². Thus, late-onset or emergent personality disorders are possible. Alternatively, individuals with lifelong personality dysfunction can have an affective denouement; they may be more likely to develop depression in old age, as some data suggest^{19,20}.

A recent body of contributions from psychoanalytic theorists and clinicians has been leading to a developmental theory of the second half of life²³. Freud's early ideas about the declining plasticity of the personality²⁴, based on experience in clinical psychoanalysis, have given way to the notion that psychosocial development is continuous throughout life. Successful disengagement from active working and parental roles, and acceptance of the inevitability of death, are several of the

proposed developmental tasks of aging. For example, awareness of one's eventual death has been thought to be marked by universal apprehension and, in some individuals, by phobias, paranoia and fear of sleep, attributed to a hypothesized "death anxiety"²⁵. Jacques²⁶ has proposed that characterologically healthier individuals have mastered the anxiety associated with awareness of death at a relatively earlier age than those with personality dysfunction, who might be blocked from acknowledging the inevitability of death until overwhelmed by it. Recently, there has been a renewed interest in psychotherapeutic work with the elderly, most proponents of which argue that change and growth is possible²⁵⁻²⁷. In the view of Neugarten²⁸ and Costa^{29,30}, personality is the critical factor in adaptation to old age.

ASSESSING THE AGING PERSONALITY

Overview of Methodological Issues

Assessing personality in the elderly is a daunting task. Each of the three basic approaches cited above carries not only its own set of assumptions but its own limitations as well. The dimensional scales have restricted clinical relevance and yield abstract, somewhat reductionistic information about the individual. On the other hand, categorical diagnoses have been criticized as culture-bound and arbitrary³, failing to "cut nature at its joints". DSM criteria, in particular, may be age-biased. Axis II criteria frequently appear to be addressing the concerns of a modal young adult, one who is expected to be establishing career and life-partner choices. If personality disorders "become less obvious in middle or old age", this may occur because Axis II does not relevantly assess the present-day experiences and behaviors of aging persons. Age bias could easily result in an underestimation of pathology, whereby symptoms are dismissed as "normative" for age. Overestimation of pathology is also possible, for example with dependency phenomena, where the realistic needs of an older person might be inappropriately viewed as symptomatic.

Psychoanalytic approaches have contributed to the study of personality and aging by encouraging the formulation of developmental theories for the second half of life. However, personality investigation of this type is focused on the individual patient, and its validity is ultimately predicated on a thorough knowledge of a few individuals; McHugh and Slavney have termed this "meaningful construct" validity³¹. While such models may explain how a patient's unique vulnerabilities and life circumstances interact to produce symptoms, large-scale empirical replication is impossible.

State-Trait Problems and Co-morbidity

Whichever theoretical-methodological model is used, the finding of age effects in personality study is frequently subject to the suspicion that they are not in fact true age effects, but rather reflect dysthymic, post-depressive or organic contaminants. This is the state-trait confound, a term that usually refers to the exaggerated self-report of some personality traits owing to the depressed state^{32,33}. For example, recovered elderly depressives have been found to have more lifetime personality dysfunction than other elderly subjects¹⁹, and in another study, twice as many recovered elderly depressives met full criteria for DSM-III-R personality diagnoses than did normal controls²⁰. While recovery from depression in these studies was carefully documented, and no personality testing was carried out during symptomatic periods, it is impossible to completely rule out depressive

influences in clinical personality assessment. The problem is especially evident in geriatric populations, in which some forms of chronic depression seem to be closely related to personality disorders. Dysthymia, a low-grade depressive syndrome appearing in as much as 15% of the geriatric population^{34,35}, could equally well be deemed an affective disorder with prominent character pathology, or a personality disorder with secondary affective symptomatology; there may in fact be subgroups of each, defined by response to antidepressant drugs³⁶.

Another chronic depressive disorder often seen in the geriatric population, also associated with significant personality psychopathology, is "double depression". Post³⁷ used the term "depressive invalidism" to describe "double depression" in the elderly, referring to a group of geriatric patients having severe recurrent depression with incomplete remissions. Finally, the term "masked depression" has been used to describe a depression syndrome, believed to be more common after mid-life, in which cognitive or somatic symptoms are more prominent than sadness, tearfulness, or other affective manifestations³⁸. Masked depression may not only present as a mixed personality disorder in geriatric patients, but has also been associated with a number of dysfunctional premorbid personality traits^{39,40}.

In addition to these chronic depressive entities, anxiety states and disorders also have considerable co-morbidity with personality disorders and have been shown to strongly influence personality assessment^{41,42}. Co-morbidity of personality dysfunction and Axis I disorders covers a wide range in geriatric patients and, as with younger patients, the relationship may take different forms. Personality disorders may variously predispose to Axis I disorders, may represent subclinical prodromes of Axis I disorders, may have a pathoplastic or interactional effect with depression or schizophrenia, or may appear as a complication or "scar" of an Axis I condition⁴³.

The Problem of Organicity

Behavioral change can occur in the context of organic brain pathology or with the use of a variety of medications, situations common in geriatric populations. However, it is not clear whether such behavior should be regarded as dysfunctional personality change or organic pathology. The approach of DSM-III-R was to categorize demented patients' symptoms concerned with the quality of affect and its regulation as an "Organic Personality Syndrome", while reserving the more purely cognitive symptomatology for the diagnosis of dementia. Spitzer *et al.*⁴⁴ proposed in the work-group stage of the development of DSM-IV that the term "organic" be eliminated. In this proposal, dementia, delirium, and amnesic disorder would be listed together in a "Cognitive Impairment Disorders" category, while the other "organic" disorders, including personality, mood and anxiety, would be viewed as "secondary". This proposal was adopted, in principle if not in detail, by the final DSM-IV⁶³, with its use of category "Personality Change Due To . . .". The nosological reclassification of organic personality disorder to secondary personality disorder emphasizes the personality aspect by listing this entity according to phenomenology rather than etiology. Implicit in the concept of "secondary" personality phenomena is an acknowledgement that personality is subject to change caused by exogenous factors or brain pathology. However, a secondary personality disorder might still be relatively enduring and pervasive, have an interpersonal focus, and lead to significant behavioral impairment; all general criteria for the diagnosis of personality disorder would be met, except for that of early onset.

Self-report vs. Informant Data

Another technical difficulty in personality study common to all theoretical viewpoints is the discrepancy between self-report and informant data²¹. For this reason, reliance upon self-report may be a particularly flawed clinical or research strategy. Among some elderly persons, including control subjects for psychiatric research, an attitude of unrealistic optimism has been observed⁴⁵. This tendency to recast the past in positive terms can be viewed either as a pathological refusal to acknowledge earlier disappointments and failures, or as a healthier attempt to reintegrate and reconcile the same experiences⁴⁶. The state influences of depression, as noted earlier, would argue in favor of informant corroboration of self-report personality data. Similarly, even the subtle cognitive impairment of early dementing illness would seriously reduce the value of self-report personality data. But who should be the informant? Adults who knew an elderly subject as a young child may no longer be alive, and adult children have not necessarily witnessed their parents' young adulthood; even then, their childhood memories might present a distorted picture. Probably it is necessary to have multiple informants, including, where possible, siblings and contemporaries of lifelong acquaintance, in order to provide the most meaningful long-range information about personality functioning.

Time-frame Considerations

Since personality is by definition concerned with established traits, or relatively enduring aspects of motivational or interpersonal behavior, over what periods of time should changes persist in order to be deemed new traits? In the clinical context, all versions of the DSM are vague on this point, stipulating that criteria for personality disorders should reflect current (past year) and long-term functioning, but not clearly requiring that symptoms be present continuously from adolescence to old age in order to qualify for a personality disorder diagnosis⁷. Also, there is no real provision for past personality disorders, those that have been present throughout much of adult life but are attenuated in old age, or for late-onset personality disorders, those first appearing or meeting full criteria in middle age or later. Recently developed standardized instruments for Axis II diagnoses may eventually produce better information on the natural course of personality disorders and the prevalence of past and late-onset disorders⁴⁶. For the moment, it is left to the clinician to select an appropriate time frame for making a personality disorder diagnosis and to determine whether current behavior reflects personality functioning, affective state, organic brain changes, adjustment reaction to age-related life events, or a combination of these factors⁴⁷.

Outcome Measures

Assuming that a suitable set of dimensions, categories or developmental milestones can be selected for personality investigation in the elderly, against what outcomes might they be expected to co-vary? Satisfaction with life, a sense of global well-being and health concerns have seemed to some authors to be the most important outcome dimensions to evaluate in older people, because they are indicators of the quality of life⁴⁸⁻⁵⁰. Longitudinal investigation suggests that, overall, well-being tends to be stable over time, probably because the frequency and intensity of both positive and negative emotions decline with age⁴⁸. However, associations between specific

personality dimensions or categories and life satisfaction in old age have not been established. At the other end of the adjustment spectrum, it has already been noted that depressed elderly patients have had abundant lifetime personality psychopathology^{19,20}. In a sample of completed suicides over age 60 years, 45% were estimated to have an Axis II disorder based on interviews with close relatives⁵¹. Nevertheless, it is not always intuitively clear what constitutes adaptive or maladaptive personality functioning in old age. For example, a relatively greater degree of risk-taking or sensation-seeking traits could be viewed as adaptive in a 20-year-old than in an elderly person; the older individual should optimally have some sensation-seeking behavior but could obviously be placed in situations of unreasonable risk without some reduction of sensation-seeking intensity. Interestingly, recovered elderly depressives have been found to have a greater preference for environmental stimulation, as suggested by higher scores on the Sensation-Seeking Scale^{52,53} than normal elderly controls⁵³, implying a complex interaction among aging, trait preference for environmental stimulation, and affective illness.

Assessment of DSM Criteria in Geriatric Patients: Clinical Notes

The diagnosis of personality disorders requires evidence of symptomatic behavior, both currently and in the past; when examining the current period of old age for evidence of personality disorder criteria, it is necessary to interpret those criteria in a geriatric context. For example, in the odd–eccentric (Cluster A) personality disorders, including paranoid, schizoid and schizotypal personality disorders, suspiciousness of exploitation or harm (a criterion of paranoid personality disorder) or social and sexual isolation (a criterion of schizoid personality disorder) should be carefully weighed against realistic dangers and social limitations faced by the older person. For this, an informant who knows the patient well is often required.

Cluster B, the dramatic–emotional personality disorders (antisocial, borderline, histrionic and narcissistic), are assigned to geriatric patients especially infrequently, possibly because they are truly “immature” syndromes that prove unstable over long periods of time, and probably also because much latitude is necessary to fit the criteria to the experiences of geriatric patients. For example, evaluation for the borderline criterion of “idealization” should include consideration of such behavior when directed toward caregivers in a hospital or nursing home. Similarly, “self-damaging impulsiveness” need not be limited to reckless driving or sexual activity, as suggested by DSM-III-R, and “frantic efforts to avoid real or imagined abandonment” should not be confused with reactions to age-related losses or institutionalization. Thus, criteria such as “self-damaging impulsiveness” might be broadened to include the geriatric context, while other criteria, such as “frantic efforts to avoid abandonment”, should be narrowed to account for the impact of realistic events.

Interpretive flexibility is also needed to assign Cluster C personality disorder diagnoses to elderly patients. Reactions and attitudes toward caregivers provide an appropriate context for many elderly patients in which to interpret Cluster C criteria, particularly those for passive–aggressive and dependent disorders. However, medical illness can transiently exacerbate dependency phenomena. For all of these disorders, it seems most appropriate to err on the side of conservatism or specificity in diagnosis. Obviously, the clinician can be most confident of the assessment when antecedents of criteria are traceable to the past, but there remains an obligation to determine what is or is not pathological in the older person’s present reality.

TREATMENT OF ELDERLY PATIENTS WITH PERSONALITY DISORDERS

There is virtually no research data to guide the comprehensive treatment of elderly patients with personality disorders. However, based upon classic and recent writings in this area^{23,25–27} and treatment reports in younger age groups, it is possible to set forth some general principles of management.

Co-morbid Axis I Disorders

It has already been noted that the depressed state may influence the assessment of personality disorders. It may also be true that, especially in the elderly, personality disorders exert their most clinically meaningful effects in the setting of major depressive disorder. For example, earlier onset of depression as well as greater severity and frequency of depressive episodes, have been found in elderly patients with personality disorders^{2,38}. Personality disorder symptoms also appear to affect elderly patients’ attitudes and behaviors toward antidepressant treatment. Personality-disordered patients have difficulty in establishing positive therapeutic alliances with psychotherapists and general physicians, a factor which in turn promotes poor adherence to either psychopharmacological or psychotherapeutic treatments. Finally, while personality dysfunction may not lengthen the depressive episode itself, in interaction with residual depressive symptoms, it appears to prolong depression-related declines in global functioning and quality of life⁵⁹.

Thus, it may be reasonable to address co-morbid Axis I conditions, particularly affective or anxiety disorders, before planning any treatment efforts directed specifically toward the personality component. In conditions such as dysthymia, where personality dysfunction and affective symptomatology at times seem to merge, it may also prove useful to defer long-term treatment planning and psychosocial decision-making until somatic treatments have reached maximum efficacy. Likewise, medical conditions that might affect the course of personality symptomatology should be fully evaluated.

Pharmacotherapy

Pharmacotherapy for geriatric personality disorder patients has not been investigated *per se* and remains an underdeveloped area. In younger age groups, there exists an older literature for borderline personality disorder, in which psychotic-spectrum symptomatology was shown to be more effectively treated by neuroleptic medication than placebo⁵⁴, and amitriptyline was usually more effective than placebo but no more effective than haloperidol in antidepressant effect⁵⁵. MAO inhibitors⁵⁶, lithium⁵⁷ and carbamazepine⁵⁸ were proposed to have limited usefulness for specific traits or symptoms. More recently, there have been some studies in mixed-age populations showing modest effects of particular medications or medication classes on the symptomatology of the three Axis II clusters. For example, haloperidol and thiothixene have been found to be useful in the treatment of patients with schizotypal or other Cluster A personality disorders, especially the transient paranoia, agitation or rages associated with these disorders⁶⁰; the selective serotonin reuptake inhibitors (SSRIs), fluoxetine and sertraline have shown some promise in treating impulsive, self-injurious and depressive behaviors associated with Cluster B personality disorders⁶¹, and fluoxetine and MAOIs have been found to decrease the kind of avoidant behaviors found in individuals with Cluster C personality disorders⁶².

Psychotherapy

Unless there is a co-morbid Axis I disorder that requires psychopharmacological intervention, the treatment of geriatric personality disorders is largely a psychotherapeutic endeavor. As described by Sadavoy²⁷, a major difference between the psychotherapy of older and younger patients is the time frame covered. Transference issues continue to be directed from childhood sources and early⁸ parental relationships, but they will often contain an overlay from experience later in life. Usually the initial focus should be on the patient's present reality and presenting problems, secondarily on historical material and on the relationship with the therapist. Then, as it unfolds, more time and effort can be devoted to clarifying and perhaps analyzing distortions in the patient's attitudes and behaviors toward the therapist. Some patients, not necessarily the highest-functioning ones, will be able to focus usefully on the relationship with the therapist to a greater degree than others. In either case, special attention must be paid to the potential for transference issues to become painful and paralyzing for both patient and therapist. Ideally, discussion of the therapeutic relationship can provide a point of shared reality, which patient and therapist can examine together. Issues from the past then emerge more naturally, in an unforced and relevant fashion. Patients probably do have a need to mourn past losses, a process that can be fostered in psychotherapy, but it should be remembered that such mourning is not done globally or in a predictable sequence²⁷.

Another factor complicating psychotherapy in the elderly is that overall treatment plans often involve family, caregivers or institutional representatives. Important persons in the patient's social sphere must be engaged in treatment strategies because of the interpersonal field in which personality psychopathology, especially the Cluster B disorders, is expressed; also, older people with disabilities may function less autonomously. This creates potential boundary problems for the therapist, as well as concerns about confidentiality; these must be spelled out to the patient and an understanding reached between the patient and therapist on exactly what information may be transmitted to others and under what circumstances this will be done. Once this is accomplished, creative use can be made of family and institutional supports.

Whatever the approach taken in psychotherapy, limited and realistic goals should be set, based upon a collaterally informed picture of the patient's long-term functioning. A psychotherapy relationship cannot reasonably be expected to resolve the psychological deficits, and the consequences of those deficits of the elderly personality disorder patient—a lifetime of failed relationships, missed opportunities and unused talents. Nevertheless, it can be hoped that for some individuals, the loss of narcissistic gratifications associated with physical beauty and vocational competence can actually foster psychotherapeutic work. Patients may find themselves in old age to be motivated for self-examination as never before. The impetus provided by aging and the press of reality may render the older patient amenable to a process of growth and change.

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Part J

Mental and Behavioural Disorders due to Psychoactive Substances

Alcohol Abuse in the Elderly

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Alcohol abuse in the elderly may be defined as the persistent and intended use of ethyl alcohol despite the problems caused by its use^{1,2}. In the elderly, alcohol abuse usually presents clinically as self-neglect, falls, confusion, lability, depression, unusual behavior, injuries, diarrhea, malnutrition, myopathy, incontinence or hypothermia^{3,4}. In fact, the elderly person with alcohol abuse problems may be hospitalized for any one of these problems. During the course of the hospitalization, one may uncover signs of characteristic addictive use of ethyl alcohol, with: (a) tolerance; (b) withdrawal symptoms; (c) loss of control of use; (d) social decline; and (e) mental and physical decline⁵. Alcohol abuse in the elderly is an under-recognized problem that has become increasingly important due to the growing numbers of elderly people⁶⁻¹⁰. Prevalence estimates of alcohol-related problems are in the range 1-6% in community-dwelling elderly, 7-22% in medically hospitalized elderly, and 28-44% in elderly psychiatric inpatients^{1,4}. In one study of older primary care outpatients, 15% of men and 12% of women regularly drank in excess of limits recommended by the National Institute of Alcohol Abuse and Alcoholism (> 7 drinks/week for women and > 14 drinks/week for men)¹¹.

EFFECTS OF ALCOHOL IN THE ELDERLY

Older people are at greater risk and more vulnerable to the toxic effects of alcohol for three main reasons:

1. A smaller volume of alcohol is required to produce the same effects as in a younger person. The elderly have a decreased volume of distribution due to decreased muscle mass, a greater proportion of fat and a smaller water compartment. These all result in a higher blood alcohol level than in a younger adult for the same amount of alcohol consumed¹²⁻¹⁶. This suggests that the elderly person's brain, liver, cardiac and other organ systems are subjected to a greater toxic effect from a given amount of alcohol. In a younger person, larger amounts of alcohol consumption may be necessary before detrimental effects from alcohol abuse become grossly evident. An elderly person may reach this threshold for hazardous use of alcohol after drinking a relatively small amount¹⁷.
2. The general decrease in the capacity to withstand stress and maintain homeostasis, as well as a higher risk for medical illness and disability in elderly people, can magnify the effects of alcohol abuse in the elderly.
3. Some organ systems may be especially susceptible to alcohol in the elderly. For example, the central nervous system appears to be more sensitive to alcohol in the older person¹⁸⁻²¹, and bone fractures are much more frequent among elderly who use alcohol than in those who do not²².

The interaction of these three main factors places the elderly alcohol-using person at greater risk for multiple impairments resulting from the use of alcohol.

There are many possible detrimental effects from alcohol abuse in the elderly. Among them are the following:

1. Driving ability can be adversely affected with the consumption of minimal amounts of alcohol. Relatively small amounts of alcohol can lead to confusion, visuospatial impairment, problem-solving deficits and motor impairment in the elderly²³⁻²⁵, which can inhibit the continuous attention and quick responses needed for driving. If the elderly person also has cognitive or sensory deficits, then the additional insults from alcohol use may make driving considerably more dangerous^{17,20}.
2. Cognitive impairments suggesting dementia may be caused by alcohol abuse^{26,27}. Although some cognitive impairment can result from even social drinking, chronic alcohol consumption has been shown to cause marked cognitive deficits, with associated cortical atrophy and ventricular dilatation on brain scan^{19,28,29}. Some researchers have suggested that alcoholism contributes to accelerated mental aging, but this is still controversial^{19,30}.
3. Elderly alcoholics have a higher prevalence of alcohol-related medical conditions than the elderly population at large. Such conditions include alcoholic liver disease, alcoholic cardiomyopathy, hypertension, chronic obstructive pulmonary disease, neurologic diseases (including cognitive brain syndromes and peripheral neuropathy), malnutrition, osteopenia, psoriasis, peptic ulcer disease and various cancers^{22,31-36}.
4. Alcohol use can adversely affect the elimination of some drugs and add to the toxicity of others. This places an elderly person with medical illness or disability who is taking prescription medication at great risk for having subtherapeutic or adverse effects from the medication^{37,38}. The magnitude of this problem is evident when one considers that the elderly receive 25% of all drugs prescribed in the USA, while comprising only approximately 12% of the population^{6,39,40,41}.
5. The depressant effects of alcohol on the central nervous system may mimic or contribute to depression in the elderly^{42,43}. Some elderly with depressed mood may resort to drinking in order to "self-medicate" themselves. This may alleviate the depressive symptoms initially, but later lead to an increase in depression, anxiety, sleep disturbances and impotence^{40,44}.
6. Alcohol can contribute to malnutrition in the elderly. Malnutrition can result from the interaction of the following factors^{45,46}:
 - (a) Food intake can be hindered if the elderly alcoholic develops depressed mood, becomes apathetic and experiences

loss of appetite. If the elderly alcoholic's impaired ambulation results in a reduced capacity to obtain food, or if limited financial resources are used to purchase alcohol instead of food, dietary intake may be restricted further.

(b) The effect of alcohol on the gastrointestinal tract is to produce malabsorption of fats, fat-soluble vitamins, calcium, magnesium, iron and zinc. The active transport of B vitamins is also impaired.

(c) Alcohol can contribute to increased losses of magnesium, phosphate, potassium and zinc through the urine. If vomiting and diarrhea occur, there may be increased loss of sodium, potassium and chloride.

(d) Alcohol use increases the requirements for folate and pyridoxine.

7. Alcohol use contributes to accidents and injuries that may lead to fractures or subdural hematomas. A study of accidental drowning in Denmark found that between one-third and one-half of adult drownings were related to alcohol intake⁴⁷.
8. Alcohol use disorder has been associated with higher mortality in a study of older public housing residents⁴⁸.
9. Alcoholism can disrupt the elderly alcoholic's family structure and cohesiveness and may even lead to family violence. This can result in dysfunctional family relationships, with consequent increased difficulty in treatment of the alcohol-related problems.

Despite the many unfavorable effects of alcohol abuse in the elderly, researchers have also reported positive aspects of alcohol use. Moderate alcohol consumption has been associated with a decreased risk of ischemic stroke in elderly subjects⁴⁹. A study of elderly Australians found that alcohol intake was associated with a significant increase in life expectancy⁵⁰.

CHARACTERISTICS OF ELDERLY ALCOHOL ABUSERS

Elderly alcohol abusers differ from younger alcohol abusers in a number of ways. Alcohol abuse in the elderly is often associated with a clustering of events, which are common in late life. These include such occurrences as job retirement, widowhood, the deaths of close friends and relatives, more medical illness and disability in oneself and one's peers, and perceived loss of a meaningful role or function. Some authors consider late-onset alcoholism to be "reactive alcoholism", where the dependence on alcohol is initiated by a need to alleviate the stresses of undergoing multiple losses. However, the extent to which alcohol abuse in the elderly is precipitated by stress from these losses is unclear. Some researchers have found little change in alcohol consumption or drinking behavior due to life stressors^{51,52}.

The time of onset of alcohol abuse may also significantly differentiate the younger alcoholic from the older one. Early-onset alcoholics have a greater amount of psychopathology and family history of alcoholism than late-onset alcoholics. Early-onset alcoholics are characterized by being male relatives of alcoholic men with histories of violence with and without alcohol, legal problems due to alcohol use, and illegal substance abuse. Late-onset alcoholics are characterized by having isolated alcohol-induced problems with health, marital relationships or self-care, and much reduced histories of arrests, violence or other substance abuse. Many elderly people with alcohol problems fall into the late-onset alcoholic group. These findings suggest that the etiology and predisposition of a person to an alcohol use disorder may differ by onset age. If this is so, the treatments and interventions for an alcohol use disorder may also differ with age of onset and need to be individualized accordingly^{53,54}.

Individual feelings towards alcohol use are affected by the cultural and historical attitudes one grows up with. For example, the experience of the American elderly alcoholic may differ from that of younger alcoholics in that the elderly alcoholic and his peers may have been exposed to the turmoil of the Prohibition era. The moral issues highlighted in this historical period may influence the willingness that some elderly may have in recognizing and accepting a diagnosis of, and treatment for, alcoholism.

THE RECOGNITION OF ALCOHOL ABUSE IN THE ELDERLY

Alcohol abuse in the elderly often comes to the attention of health professionals through presentation with a non-specific medical or psychiatric symptom, such as self-neglect, falls, confusion, lability, depression, unusual behavior, injuries, diarrhea, malnutrition, myopathy, incontinence or hypothermia. In cases where alcohol abuse is suspected, alcohol dependency must be considered. Alcohol dependency is suggested when there are: (a) tolerance; (b) withdrawal symptoms; (c) loss of control of use; (d) social decline; and (e) mental and physical decline.

Tolerance to alcohol may be assessed by establishing a reliable history of the patient's drinking pattern. Corroboration from family members and others close to the patient may be crucial. Tolerance is suggested if the patient exhibits a quantity and frequency of drinking which is increased over his baseline pattern of drinking. A patient with tolerance to alcohol will require a greater quantity of alcohol to achieve the same amount of inebriation that a lower quantity had been able to achieve previously. Tolerance is strongly suggested if there has been at least a 50% increase in the amount of alcohol required to attain a given effect, a blood alcohol level of 150 mg without intoxication, or the equivalent use of one-fifth gallon (750 mg) of alcohol or more in 1 day by a 180 pound person⁵⁵.

Withdrawal symptoms occur when a patient who is tolerant to alcohol experiences a rapid decrease in blood alcohol concentration. Symptoms of the alcohol withdrawal syndrome include tachycardia, with a pulse of greater than 110 beats/min, tachypnea, hypertension, low-grade fever, sweating, nausea, vomiting, hand tremors and increased anxiety. In some cases, the patient may develop seizures or delirium tremens with confusion, agitation and visual hallucinations. An elderly patient undergoing withdrawal may experience one or all of these symptoms^{55,56}.

Loss of control means that the patient is no longer able consistently to choose the amount of alcohol he/she will consume in a given situation. He/she may also experience blackouts and behave and feel in unpredictable ways⁵⁵.

Social decline in the elderly alcoholic is assessed from a baseline of age-appropriate behaviors⁵⁵. Many elderly people no longer hold a steady job, do not drive or hold a driver's license, and have lost many of their close friends and associates with whom they used to socialize. Thus, it may not be as appropriate to assess for social decline by investigating these areas of the elderly patient's life as it would be in a younger patient. However, it is relevant and revealing to ask elderly people whether they are in contact with their children or grandchildren, and to what extent. It is also useful to find out whether the patient's relatives express any concern about the patient's alcohol use. Investigating the patient's functioning with respect to his/her hobbies or other enjoyed activities can also be useful.

Physical, psychological and laboratory findings may also uncover problems with alcohol use⁵⁵. Addictive alcohol use can lead to malnutrition, gastrointestinal upset and bleeding, delirium, falls, depression, hypertension and neglect of self. Recurrent diseases of the stomach, pancreas or liver may also be caused by excessive alcohol abuse. These medical conditions often bring the

elderly alcoholic to clinical attention. Laboratory results of macrocytosis, elevated mean corpuscular volume, and increased liver enzyme levels, especially γ -glutamyl transpeptidase, may correlate with alcohol abuse in the elderly. Blood alcohol levels, and urine or breath tests for alcohol, may be used to confirm alcohol intoxication.

Assessment of tolerance, withdrawal, loss of control, social decline and mental and physical decline are useful clinical parameters to recognize and diagnose alcohol addiction. Several screening instruments have been devised to help clinicians recognize alcoholism. These scales typically assess the quantity and frequency of drinking, social and legal problems resulting from alcohol abuse, health problems related to excessive alcohol use, symptoms of addictive drinking, and/or self-recognition of alcohol-related problems⁵⁷⁻⁵⁹. Many of these instruments have been validated for younger populations, but not specifically for the elderly. The validity of the CAGE screen for alcoholism in the elderly has been examined empirically⁶⁰. "CAGE" is a mnemonic for the questions: Have you ever felt a need to Cut down on drinking? Have you ever felt Annoyed by others inquiring about your drinking? Have you ever felt Guilty about drinking? Do you ever use alcohol for an Eye-opener? If two or more of these questions are answered positively, a need for more extensive evaluation for alcohol abuse is indicated. Another more detailed screen is the Michigan Alcoholism Screening Test—Geriatric Version (MAST-G)⁶¹. In addition, the original MAST, scored with weighted (MAST) and unit scoring (UMAST), and two shorter versions, the Brief MAST (BMAST) and Short MAST (SMAST), have been tested in the elderly⁶². Researchers found that the MAST and UMAST gave excellent sensitivity and specificity for alcohol abuse in the study population of 52 hospitalized elderly male alcoholics, matched with 33 non-alcoholic controls. The MAST and UMAST may be useful screening instruments to help recognize alcoholism in the elderly.

THE TREATMENT OF ALCOHOL ABUSE IN THE ELDERLY

Once the diagnosis of alcohol abuse is confirmed, the first step in treatment is a thorough evaluation to identify any other coexisting medical or psychiatric problems. Treatment for these problems must be initiated at the same time as the patient is detoxified. Providing adequate nutrition and hydration is especially important in the elderly alcoholic, due to increased nutritional problems and impaired thirst mechanisms in the elderly. Benzodiazepines are generally avoided in the detoxification of elderly alcoholics, due to their potential for causing delirium. However, if significant withdrawal symptoms occur, benzodiazepines should be administered^{56,63,64}. Elderly patients may experience more severe withdrawals from alcohol and require higher doses of benzodiazepines than younger patients. In general, detoxification with benzodiazepines with a short half-life is recommended in the elderly. However, these medications may not provide adequate anticonvulsant effect and the use of longer-acting benzodiazepines may be necessary⁵⁶.

Once the elderly alcoholic patient is detoxified, adequate relapse prevention and rehabilitative treatment is crucial for the patient to maintain sobriety. The elderly alcoholic must first come to an acceptance of his/her alcohol abuse problem. Family members or others who are close to the patient may be able to help the patient break through the denial regarding alcohol abuse seen in many alcohol abusers. Family members may also be instrumental in motivating the patient to stop drinking. Patient and family education about the effects of alcohol must be provided, and the need to abstain from alcohol must be stressed. A behavioral and self-management treatment module has resulted in marked success

in treating alcohol use problems in the elderly⁶⁵. This module educates the elderly alcoholic about drinking behavior, the acquisition of self-management skills, and the re-establishment of social networks.

The elderly alcoholic patient's recovery and rehabilitation is an ongoing process. The patient needs to learn to readjust to life without alcohol. The patient's family and close relatives and friends can help him/her in this endeavor by supporting sobriety in the patient and incorporating the non-drinking patient into their lives without the presence of alcohol. Family members may have been "enabling" the patient to drink, and they need to be made aware of these patterns of behavior and change them through family education and family therapy. If the elderly alcohol abuser is a parent and the enablers are his children, the role reversal that is inherent in the children's setting limits on their parents may make this task especially difficult. Group therapy may help the patient adjust to a non-alcoholic lifestyle. In this setting, he/she can develop non-alcohol-related social skills and learn to bond with others in safe surroundings, free of the context of alcohol. If elderly alcoholics can be treated in age-specific groups, they may remain in treatment significantly longer and be more likely to complete treatment than those treated in mixed-age groups⁶⁵. Involvement in Alcoholics Anonymous with people with whom the patient feels comfortable and can consider his/her peers is important, especially if he/she has no close relatives or friends. The relationships with others that can be formed in these settings can provide a replacement for the alcohol to which the patient was bonded previously. Family involvement with the patient in Alcoholics Anonymous and in other affiliated groups such as Alanon, Alateen or Alatot is also important, as alcoholism adversely affects family members who also need support, education and treatment.

Use of alcohol-deterrent medications, such as naltrexone or disulfiram, may serve as adjunctive treatments. These drugs may help motivated alcohol abusers to reduce the quantity of alcohol used or the number of drinking days. However, these medications generally are not effective unless prescribed and monitored as part of an overall, multidisciplinary treatment and relapse prevention plan. Opioid receptor antagonists, such as naltrexone or related compounds, may be less toxic than disulfiram in elderly patients⁶⁶⁻⁶⁸.

The cultural aspects of alcohol abuse intervention are also worthy of consideration. The above recommendations for treatment are a description of treatment conceptualizations in the USA. These interventions may need to be modified for other countries. In instituting any type of treatment or intervention, it is important to consider the context of the problem being addressed and the context into which the treatment or intervention will be instituted. The consumption of alcohol can take on a variety of cultural meanings. To some groups, ethanol is a food or is associated with religious rituals. For others, it is a means to relax and calm one's nerves. For still others, alcohol is considered to be a sinful intoxicant used by those of weak moral fiber. To intervene most effectively with an aged person for whom alcohol consumption has become a problem, it is important to understand what meaning the use of the alcohol has for him as an individual, a family member and as part of the greater society, including his cultural group. The contextual meaning of the change in alcohol use must likewise be considered. To understand these contextual meanings most effectively, the clinician must be aware of what his/her inherent assumptions may be about these contextual meanings and try not to confound with his/her own biases his/her understanding of the situation. Members of the patient's contextual and cultural groups may be very helpful in providing meaningful insights into these understandings. Once these cultural factors are understood, interventions to reduce the problematic alcohol consumption to the desired outcome of abstinence or

perhaps of less harmful alcohol use, can be creatively formulated and instituted more effectively and efficiently.

CONCLUSION

In summary, alcohol abuse in the elderly is a significant problem that needs to be addressed aggressively. Recognition and diagnosis of alcohol abuse may be more difficult in the elderly. The biological and psychosocial losses and decline that are often used to identify alcoholism in younger people can occur in many elderly who do not abuse alcohol. The clinical presentation of an elderly alcoholic is often with a medical condition that may be masked by other medical diagnoses. For these reasons, the clinician must be especially alert to the possible presence of alcohol abuse in this age group. If alcohol abuse is identified, the clinician must then be prepared to direct or initiate treatment in a culturally appropriate, sensitive, flexible and creative manner that is geared to the elderly individual and his/her family and significant others.

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Epidemiology of Alcohol Problems and Drinking Patterns

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A key finding from the Epidemiologic Catchment Area (ECA) community studies of adults 18 or older conducted in the USA two decades ago was the high prevalence of alcohol use and dependence, as defined by DSM-III criteria^{1,2}. The lifetime prevalence of alcohol abuse/dependence in the ECA sample was 13.8%. Gender differences were significant. Among men, the lifetime prevalence was 23.8%, while among women the prevalence was only 4.6%. Age differences in lifetime prevalence were noted for both males and females. The lifetime prevalence of alcohol abuse/dependence for men in the ECA was lowest in those 65 or older (14%) compared to 21% in those ages 45–64, 28% in those ages 30–44 and 27% in those ages 18–29. The 1 month prevalence among men 65 or older was 1.93%. A similar decreasing prevalence with age was seen in women, with a lifetime prevalence of 1.49% in women 65 or older and a 1 month prevalence of less than 1% (0.4%).

While the proportion of elderly with defined alcohol abuse or dependence may appear low, alcohol consumption at any level may potentially be problematic in older adults. Because of decreased lean body mass and smaller volume of distribution, higher peak ethanol concentrations per dose are found in older compared to younger subjects³. Also, even in small amounts, alcohol may exacerbate or mask symptoms of illness. Finally, many elderly are users of both prescription and over-the-counter medications that may interact with alcohol⁴. As the numbers of elderly will increase significantly over the next several decades (as the post-World War II generation ages) and alcohol use rates are not likely to decrease, alcohol use in older adults is an important public health concern. In addition, females are increasing their rate of alcohol use and make up a larger proportion of the elderly, potentially resulting in increased prevalence overall⁵.

Recent reviews have focused on the epidemiology of alcohol use among older adults^{6–11}. These reviewers have concluded that alcohol use in older adults in excess of recommended limits is an important clinical issue. Chermack *et al.*¹² examined the relationship between alcohol consumption patterns and the presence of DSM-III-R alcohol symptoms among 443 current drinkers, 55 years of age or older, and found that both average daily consumption and days of heavy drinking in the past year independently predicted symptom status. Consumption levels for men and women were only different for problem drinkers. The authors suggest that their results support the recommendation that moderate consumption levels should be lower for older than for younger adults, with a recommendation not to exceed one drink/day.

The prevalence of any alcohol use and the prevalence of heavy drinking among elders, as well as the factors and outcomes associated with drinking, are addressed below.

PREVALENCE OF ALCOHOL USE

The prevalence of alcohol use has been shown in cross-sectional studies to decline with age^{13–17}. Early studies in the USA noted this age difference^{13,14}. In 1968, Cahalan and Cisin¹³ conducted a national survey of drinking practices in the USA using a representative sample of 2746 persons aged 21+, and reported the prevalence of alcohol use declined with age. Among adults aged 60+, 65% of the males and 44% of the females drank alcohol within the past year. The prevalence was lower than that observed in younger males (84% in men aged 21–29 and 86% in men 30–39) and in younger females (70% in women 21–29 and 72% in women 30–39). In a probability study of adults living in western New York, conducted 25 years ago, the prevalence of alcohol use was lower in persons aged 60 or older than in those under 60, and the proportion of adults with alcohol-related problems was much less in those 60+. The proportion of moderate and heavy drinkers was also lower among those aged 60+ than in those younger. Only 7% of those aged 60+ were heavy drinkers, compared to 24% of those aged 50–59 and 30% of those aged 18–49¹⁴.

Recent studies have confirmed these earlier findings. The prevalence of alcohol use in the past year among persons aged 55+ participating in the National Longitudinal Alcohol Epidemiologic Survey was 29.5%, and there was an inverse relationship between alcohol use and increasing age¹⁵. In the 1988 Alcohol Supplement of the National Health Interview Survey (NHIS), the prevalence of alcohol use, defined as 12 or more drinks in the past year, was 60.9% in those aged 18–29, 58.3% in those 30–44, 47.3% in those 45–64 and 30.4% in those 65+. The prevalence of lifetime abstainers in those aged 65+ was 30.8%, compared to only 17.9% in those 45–64 and 13.1% in those 30–44. The prevalence of lifetime infrequent drinking and former drinking also increased with age¹⁶. Ruchlin used data from the 1990 Health Promotion and Disease Prevention supplement to the National Health Interview Survey to examine prevalence of alcohol consumption in adults aged 55+. A total of 46% reported that they had consumed alcohol during the past year. The study found a continuous decline across age groups, from 52.9% of those aged 55–64 categorized as current drinkers within the past year, compared to 24.7% of those aged 85+. Of the sample aged

55+, 17% reported that they had consumed alcohol every day in the past 2 weeks. In controlled regression analyses, more people aged 65–74 drank every day, compared to those 55–64 (OR = 1.36), but people aged 75+ drank less than those 55–64. Males and Whites used alcohol more frequently and were more likely to be heavy drinkers than females and non-Whites. The lower one's perceived health status, the lower the odds of drinking every day. Believing excessive drinking increases the chances of getting cirrhosis of the liver decreased the odds of moderate and heavy drinking¹⁷.

There are various explanations why the prevalence of alcohol use is lower in the elderly. Selective survival may be a factor, in that persons who drink are less likely to survive to older ages. Cohort effects are also possible. Persons who grew up in the era of Prohibition and economic depression prior to World War II may have had lower alcohol use throughout their lives⁵. Studies have also shown that some elderly decrease their use of alcohol as they grow older. Barnes found half of the subjects aged 60+ who were current abstainers reported that they were former drinkers, with "bad health" most often given as a reason for giving up drinking¹⁴. Busby *et al.*¹⁸ investigated alcohol use in a community-based sample of adults aged 70+ in New Zealand. Both frequency and quantity of intake decreased with age. A total of 60.1% of the men and 30.3% of the women said they drank less compared to middle age, and only 7.4% of the men and 11.1% of the women said they drank more. The main reasons cited for decreased use of alcohol were change in health and fewer social opportunities, with reasons cited for increased intake being more time and money¹⁸. Similarly, a significant proportion of male current drinkers, but not abusers, selected from medical admissions aged 65+ in the UK reported heavier drinking in the past. The most frequent reasons given for reduction in intake were onset of ill-health (36%), loss of social contact (28%) and financial difficulties (19%)¹⁹.

Using data from the Normative Aging Study, Glynn *et al.*²⁰ studied generational effects on alcohol consumption in a sample of 1859 male adult volunteers interviewed at baseline and 9 years later. Older men drank less than younger men at both assessments, yet there was no tendency for men to decrease their consumption levels over time. Each older cohort had a lower prevalence of problems with drinking than the next youngest cohort. The authors suggested that their results show that aging is not as important as generational changes in prevalence²⁰. In contrast, Adams *et al.* followed a cohort of 270 healthy community-dwelling elderly aged 60+ over 7 years²¹. At baseline, the investigators found a decline in the percentage of drinkers with increasing age²². In the 7 year follow-up, there was a 2%/year decline in the percentage of subjects consuming any alcohol, but mean alcohol intake did not change for those who continued to drink, except among heavy drinkers, suggesting an age-related decline rather than a cohort effect²¹. Smart and Adlaf²³ pooled data from four cross-sectional surveys of adults in Ontario and found no dramatic changes in alcohol use between 1976 and 1984 among persons aged 60+. Elderly respondents were more likely to report abstinence from drinking than those younger (40% of those aged 60+ compared to 15% of those 30–59 and 13% of those 18–29). Of the elderly respondents, 17% reported consuming five or more drinks in a single sitting, a much lower prevalence than among those 30–59 (44%) and those 18–29 (60%)²³.

Regardless of a decline with age, the prevalence of alcohol use in the elderly remains high. Meyers *et al.* found, in their study of the drinking behavior of 928 residents of Boston aged 60+, that 53% were abstainers, 26% had less than one drink/day, 16% one to two drinks/day and 6% two or more drinks/day²⁴. In a sample of 270 healthy men and women aged 65–89 living in the southwestern USA, 48% of the participants reported in their

3 day diet record they had consumed alcohol, with 66% reporting that they consumed alcohol at least monthly²².

The prevalence of alcohol use in the recently conducted multi-site Established Populations for Epidemiologic Studies of the Elderly (EPESE) studies of persons aged 65+ varied by site. In East Boston, 70.5% of the sample drank alcohol in the past year and 54.7% had used alcohol in the past month. Similar findings were reported from New Haven; 65.8% had used alcohol in the past year and 51.9% in the past month. The proportions were lower in Iowa and North Carolina. In the Iowa sample, 46.3% had used alcohol in the past year, while 31.2% had done so in the past month. In the North Carolina EPESE, 33.4% had used alcohol in the past year and 24.6% in the past month. Men were more likely to report use of alcohol in the past month than were women^{25,26}.

Samples from clinical populations have found similar proportions of high use. In one of the first studies done in the UK, Bridgewater *et al.* interviewed 101 patients aged 60+ from a general practice and found that 92% of the men and 77% of the women used alcohol²⁷. Iliffe *et al.* studied 241 patients from general practice aged 75+ and found that 51% of the men and 22% of the women reporting using alcohol in the past 3 months²⁸. Using data from the Liverpool Longitudinal Study of Continuing Health in the Community, Saunders *et al.*²⁹ reported, among 1070 men and women aged 65+ randomly selected from patient rosters, that 10.5% admitted to drinking more or less every day (17.7% men and 6.1% women). At the 3 year follow-up, one-fifth of the subjects were regular drinkers, drinking on at least one occasion per week²⁹. Among a sample of 132 general hospital patients aged 65+ in Leiden, The Netherlands, the prevalence of alcoholism was 9% (13% among men and 7% among women)³⁰. Callahan and Tierney³¹ found the prevalence of alcoholism to be 10.6% among 3954 primary care patients aged 60+. Patients with alcoholism were more likely to be younger, have fewer years of education, and be male, Black, smokers and malnourished³¹. In a sample of 539 medical admissions aged 65+ in the UK, the prevalence of alcohol abuse was 7.8%. An additional 29.7% of the sample who were neither abstainers nor occasional drinkers nor alcohol abusers drank regularly; 42.9% of the men and 15.9% of the women¹⁹.

In summary, research studies conducted in both community and clinical samples over the last several decades have consistently shown that alcohol use among older adults is not uncommon, and that over half of community-dwelling older males may consume alcohol on a regular basis. While the prevalence of use may be lower than that seen in younger populations, it is still not clear whether this is a cohort or an age effect.

PREVALENCE OF HEAVY DRINKING

Estimates of the prevalence of heavy use are also quite high in the elderly. Cahalan and Cisin reported from their community sample that 20% of the males and 2% of the females aged 60+ were classified as heavy drinkers¹³. Meyers *et al.*²⁴ found in their sample of Boston residents 60 or older that 1% were self-reported problem drinkers. All problem drinkers had long-term drinking problems and were less likely to be satisfied with their social relationships. While the proportion of problem drinkers was lower in those aged 75+ compared to those aged 60–75, the proportion of drinkers who report problems was similar for both groups²⁴. Mirand and Welte studied a sample of community-dwelling adults aged 60+ in Erie County, New York, and found the prevalence of heavy drinking was 6%³². In their sample of 270 healthy male and female volunteers aged 65+, Goodwin *et al.* found that 17% drank more than 30 g alcohol/day on average²². In the EPESE, the percentage of persons who drank two or more

ounces of absolute alcohol/day was 8.4% in East Boston, 6.6% in New Haven, 5.4% in Iowa, and 7.2% in North Carolina^{25,26}.

The prevalence of heavy drinking in clinical samples is similar. Adams *et al.* screened 5065 patients aged 60+ seen in primary care and found 15% of the men and 12% of the women regularly drank in excess of recommended limits; 9% of the men and 2% of the women reported regularly consuming more than 21 drinks per week³³. Bridgewater *et al.* reported that the prevalence of heavy drinking was 27% in men and 9% in women in their sample of 101 patients from general practice²⁷. In their Liverpool study, Saunders *et al.* found a total of 6.1% of the men and 2.4% of the year 3 subjects regularly exceeded safe consumption limits. These figures translate into 19.5% of the men and 19.6% of the women being regular drinkers who were exceeding sensible limits. They observed a decline with age in the proportion of subjects who were regular drinkers²⁹. Bristow and Clare interviewed 650 medical and geriatric admissions over 65 and found 9% of the men but few (0%) of the females drank in excess of recommended safety limits. Another 10% had cut down, primarily because of medical reasons. Compared to the non-drinkers and light drinkers, the heavy drinkers were more likely to smoke, not to be married, and to have some impairment of mobility³⁴. Iliffe *et al.* found 3.6% of the men in their sample and 3.2% of the women admitted consuming more than 21 and 14 units of alcohol per week, amounts in excess of recommended safe limits for males and females. Neither drinking status nor total weekly alcohol consumption was associated with age, cognitive impairment, depression, falls or inpatient or outpatient care²⁸.

Although the overall prevalence is low, heavy drinking among older adults is of much concern, with perhaps as much as 20% of users drinking in excess of recommended limits.

IDENTIFICATION OF PROBLEM DRINKING IN THE ELDERLY

Physicians may have difficulty recognizing alcoholism in elderly subjects. First, screening instruments used in younger populations may not be reliable for older adults. Adams *et al.* compared responses to a beverage-specific self-administered questionnaire about the quantity and frequency of alcohol use and episodes of binge drinking to the widely used CAGE questionnaire (Cut down, Annoyed by criticism, Guilty about drinking, Eye-opener drinks)³⁵ in 5065 primary care patients aged 60+ and found the CAGE performed poorly in detecting heavy or binge drinkers³³. Lutrell *et al.* similarly concluded the sensitivity of standardized screening instruments was low in patients aged 65+ admitted as emergencies³⁶. In addition, many of these screening instruments inquire about frequency and quantity of alcohol use. Many elderly may drink daily but in smaller amounts. These small quantities, however, may cause problems because of interactions with medications and chronic illness⁴.

Second, criteria for alcoholism often include problems with social and/or occupational functioning. However, many older adults are less likely to be married or employed, and therefore less likely to report marital or job problems. Older drinkers may be more likely to maintain a "low profile" and not cause public disturbances resulting in legal problems⁵.

Finally, physicians may fail to diagnose alcoholism in the elderly, perhaps because they often fail to obtain alcohol histories³⁷ or because they confuse perceived symptoms of aging with symptoms of alcoholism⁵. Curtis *et al.* screened all new admissions to the medical service at the Johns Hopkins Hospital for alcoholism using the CAGE³⁵ and the Short Michigan Alcohol Screening Test (SMAST)³⁸. The prevalence of alcoholism was 27% in patients under age 60 and 21% in patients 60+. These age differences were not significant. However, 60% of screen-positive

younger patients were identified as having alcoholism by their house officers, compared to only 37% of those aged 60+. Elderly patients with alcoholism were less likely to be diagnosed if they were White, female or had completed high school³⁹.

In a similar study, Adams *et al.* screened patients aged 65+ who came to the emergency department for alcoholism. Using their criteria of either CAGE-positive or self-reported drinking problem and alcohol use within the past year, they found the current prevalence of alcohol abuse was 14%, with a high prevalence (22%) among those presenting with gastrointestinal problems. Physicians, however, detected only 21% of current alcohol abusers⁴⁰. In a study conducted in The Netherlands, scores on the Dutch version of the Munich Alcoholism Test⁴¹ and medical records were obtained from 132 patients aged 65+ staying at University Hospital Leiden. Two-thirds of the alcoholic patients were recognized by the attending physician³⁰. Finally, medical staff identified only 33 of 99 problem drinkers among inpatients aged 65+ in three hospitals in New South Wales⁴².

These studies consistently show that older problem drinkers may be more difficult to identify as a result of poor screening instruments, failure of clinicians to consider problem drinking as a possible contributing diagnosis, and difficulty in separating problems caused by alcohol from those caused by other diseases.

FACTORS ASSOCIATED WITH ALCOHOL USE IN THE ELDERLY

Among the elderly, alcohol use has been shown to be associated with male gender^{17,22,31-33,43}, higher income²², more education^{22,33}, lower socioeconomic status³², being married³³ and current or former smoking³²⁻³⁴. Other studies have found alcohol use associated with less education³¹. Goodwin *et al.* found no differences in social support between elderly drinkers and non-drinkers and no relationship between alcohol intake and emotional status²².

Two factors often associated with alcoholism in late life are depression and impairments in cognitive functioning. Saunders *et al.*, using data from their Liverpool study, reported 44% of men 65 or older with a history of heavy drinking were given current psychiatric diagnoses, compared with 12% of the men without a history of heavy drinking. The most common diagnoses were depression and dementia. The association between drinking history and current psychiatric morbidity was not explained by current drinking habits⁴⁴. Finlayson *et al.* studied 216 patients aged 65+ admitted to the hospital for treatment of alcoholism. Patients with late-onset alcoholism (aged 60+) reported a higher frequency of life events associated with problem drinking compared to those with earlier onset. The most common comorbid psychiatric disorders were tobacco dependence (67%), organic brain syndrome (25%), atypical or mixed organic brain syndrome (19%) and affective disorder (12%); 14% of the patients had a drug abuse or dependence problem, all using legally prescribed drugs⁴⁵. Similarly, Speckens *et al.* found that alcoholics used more psychotropic drugs compared to non-alcoholics and suffered more often from organic brain disease³⁰.

Iliffe *et al.*, however, found among patients aged 75+ that current drinking was not related to age, depression or mental status score²⁸. Goodwin *et al.* found that those elderly who consumed alcohol performed better on the cognitive functioning tests, but no relationship was found between past alcohol consumption and present cognitive performance. The authors concluded that alcohol may not impair cognitive functioning in the elderly²². In a study of adults aged 70-75 in Italy, self-reported alcohol consumption was associated with male gender, better mood, less cognitive and functional impairment, better health, not living alone and being married, while CAGE-positive alcoholism

was associated with male gender, poorer cognitive function and income dissatisfaction. The groups did not overlap much, suggesting some positive aspects of alcohol use, and that self-reported consumption (they used the top 10% of quantity) may not necessarily be associated with alcoholism⁴⁶.

Mangion *et al.* found that men aged 65+ classified as alcohol abusers were more independently mobile than those not abusing alcohol, suggesting greater physical fitness¹⁹. Bristow and Clare, however, found that in an elderly inpatient sample, drinking in excess was associated with impairment in mobility³⁴.

Mirand and Welte studied the relationship between health-orientated lifestyle and heavy drinking among the elderly in Erie County, New York, and reported that the prevalence of heavy drinking was 6%. Heavy drinking was positively associated with being male, having suburban residency and currently smoking, and negatively associated with SES, rural residency and degree of health orientation. Age and level of active lifestyle were not related to drinking³².

Molgaard *et al.* found racial differences among 65+ subjects in drinking level, both before and after age 40. Among Whites, 73.8% reported drinking after age 40, compared to 48.6% of Blacks and 44.3% of Mexican-Americans. A higher proportion of Whites than Blacks or Mexican-Americans reported more minimal drinking before and after age 40. However, there were no statistically significant differences for severe drinking among the groups⁴⁷.

ONSET OF PROBLEM DRINKING IN THE ELDERLY

Recent research has focused on the age-of-onset among elderly problem drinkers. Specifically, in several studies two groups have emerged. First, there are elderly who have had problems with alcohol most of their adult life and have survived to old age, generally referred to as "early-onset problem drinkers". There are also elderly who may or may not have consumed alcohol earlier in their lives, but who do not become problem drinkers until later in their adult life. This group is generally referred to as late-onset problem drinkers and the incidence of problem drinking has been hypothesized to be a result of a stressor.

Atkinson *et al.* studied the age of onset among 132 60+ men admitted into an outpatient treatment program and found onset after age 60 in 15% of the sample and in 29% of the sample aged 65+. Later-onset alcohol problems were milder and associated with greater psychological stability. Treatment variables were better predictors of treatment outcome than age of onset⁴⁸.

Brennen and Moos reported that late middle-aged problem drinkers reported more negative life events, chronic stressors, and social resource deficits than did non-problem drinkers⁴⁹. However, in their same population, Brennen and Moos⁵⁰ studied men and women aged 55–65 and compared age-related loss events, overall negative life events and chronic stressors reported by late-onset, early-onset and non-problem drinkers. Late-onset problem drinkers consumed less alcohol than early-onset ones, reported fewer alcohol-related problems, functioned better, and had fewer stressors than early-onset drinkers. They did not find evidence for an association between age-related loss events and the onset of late-life drinking patterns. Similarly, Barnes found that neither widowhood nor retirement was related to heavy drinking. Heavy drinking was twice as prevalent among those subjects aged 60+ who were employed compared to unemployed¹⁴.

Brennen *et al.*⁵¹ found gender differences among late-middle-aged and older problem drinkers. Specifically, women with drinking problems consumed less alcohol, had fewer drinking problems, and reported more recent onset of drinking problems than did male problem drinkers. The female problem drinkers also used more psychoactive medications, were more depressed

and were less likely to seek treatment. Osterling and Berglund studied gender differences in first-time admitted alcoholics aged 60+ to a treatment center in Sweden and found that age of onset of problem drinking occurred significantly later in females compared to males. During the period 1988–1992, sex ratios indicated a significant convergence of female patients compared with a decade earlier. The authors make it clear that it is not known whether this represents an increase in problem drinking in elderly females, or whether females feel more free to seek treatment than a decade earlier⁵². Hurt *et al.*⁵³ studied 216 patients aged 65+ admitted to an alcoholism treatment program. Early-onset alcoholism was present in 59% of the men and 51% of the women, while late-onset alcoholism was present in 39% of the men and 46% of the women (time of onset was not available for 2%). Few differences were noted between the two groups⁵³.

Moos *et al.*⁵⁴ followed their cohort of problem drinkers aged 55–65 for 1 year. Remitted problem drinkers were those who did not experience any problems in the 1 year follow-up period. At baseline, the to-be-remitted problem drinkers consumed less alcohol, reported fewer drinking problems, had friends who approved less of their drinking, and were likely to seek help from mental health practitioners. In addition, late-onset problem drinkers were more likely to remit over the 1 year period.

OUTCOMES ASSOCIATED WITH PROBLEM DRINKING IN THE ELDERLY

Some medical disorders have been found to be more prevalent in elderly with a history of alcohol use or current use. Hurt *et al.*⁵³ described 216 elderly patients aged 65+ treated for alcoholism in an inpatient treatment program. The frequency of serious medical disorders among this group was higher than what would be expected for the overall population aged 65+. Hypertension was less frequent among these patients, while alcoholic liver disease, chronic obstructive pulmonary disease, peptic ulcer disease and psoriasis were more prevalent among the alcoholic group. The frequencies of ischemic heart disease, cerebrovascular disease and diabetes mellitus were about the same as would be found in the general elderly population⁵³. Bristow and Clare found, in their inpatient sample, that excess drinking was associated with more non-malignant respiratory disease and less ischemic heart disease³⁴.

Increased hospitalizations have also been linked with alcohol use in the elderly. Callahan and Tierney reported from their sample of patients aged 60+ that patients with alcoholism were more likely to be hospitalized (21.5% vs. 16.9%) ($p=0.02$) within the year following the interview, compared to those without alcoholism³¹. Using 1989 hospital claims data, the prevalence of alcohol-related hospitalizations among people aged 65+ in the USA was 54.7 per 10 000 population for men and 14.8 per 10 000 for women, a proportion of hospitalizations among the elderly similar to that seen for myocardial infarction⁵⁵.

Alcoholism has also been linked with mortality in the elderly. Callahan and Tierney found in their sample of elderly patients that those with alcoholism were more likely to die within 2 years than those without evidence of alcoholism, 10.6% compared to 6.3% ($p=0.001$), controlling for age, gender, race, education and smoking history³¹. In their study of inpatients treated for alcoholism, Hurt *et al.* followed 60 of their patients for an average of 5.2 years (range 2–11 years). A total of 32% of the alcoholic patients had died by follow-up. Of those who died, 47% of the deaths could be attributed to the patient's alcoholism⁵³. Colsher and Wallace, using data from the Iowa 65+ Rural Health Study, found that 10.4% of the men had self-reported histories of having been previously heavier drinkers. Three-year mortality was higher among this group, compared to those men without a

history of heavy drinking⁵⁶. Mellstrom *et al.* examined a cohort of 468 70-year-old Swedish men in 1971–1972, and reinterviewed them 5 years post-baseline (1976–1977) ($n=342$) when they were 75 years old. In addition, they interviewed a control group of 70-year-old men in 1976–1977. Registration at the Temperance Board (recidivists) was used to measure previous alcohol abuse or large-scale consumption. Morbidity from diabetes and chronic bronchitis was higher in the recidivists, as was the overall 5 year mortality rate. Impaired functioning and a high consumption of institutional care were also more frequent among the recidivists⁵⁷.

Other researchers have also found impairments in functioning in those elderly with a history of alcohol use. Colscher and Wallace found that men with a history of heavy drinking had more illness, poorer self-perceived health, more physician visits, more depressive symptoms, lower levels of life satisfaction and smaller social networks compared to non-heavy drinkers and non-drinkers. The authors concluded that a history of heavy drinking was predictive of impairments in physical, psychological and social health and functioning among elderly men⁵⁶.

Some studies have found a protective effect of moderate alcohol use in the elderly. Scherr *et al.*, using the EPESE data from three sites, found that low to moderate alcohol consumption was associated with lowered 5 year total mortality as well as cardiovascular mortality in two of the sites, East Boston and New Haven. In Iowa, there were no differences in mortality by alcohol consumption. There was no association with cancer mortality found at any of the three sites. Patients with a baseline history of heart attack, stroke or cancer were excluded from their analyses⁵⁸. LaCroix *et al.* found, by following three of the EPESE cohorts of elderly aged 65+ for 4 years, that risk of losing mobility, defined as the ability to climb up and down stairs and walk a half a mile, was associated with not consuming alcohol, compared with small to moderate amounts of alcohol consumption⁵⁹. Galanis *et al.* recently conducted a longitudinal study of drinking and cognitive performance in elderly Japanese-American men and found lower scores on cognitive functioning tests in non-drinkers and heavy drinkers (more than 60 ounces of alcohol/month). Compared with non-drinkers, the risk of a lower score (more errors) on a cognitive functioning test was lowered by 22–40% among men who consumed 1–60 ounces of alcohol/month⁶⁰.

Finally, in a sample of 216 inpatients treated for alcoholism, the treatment outcome was favorable (i.e. the patient was either abstinent since treatment or abstinent with three or fewer minor slips) for 28% of the cohort of 60 patients, showing that the elderly alcoholic can be successfully treated⁵³.

SUMMARY

In summary, alcohol use, including drinking in excess, among older adults is prevalent, particularly among males. Because of interactions with chronic disease and use of prescribed medications, the use of alcohol among elderly individuals is an important health concern. Based on current use, the prevalence of alcohol use in future generations may be even higher. Screening and questioning for alcohol problems should be routine, since effective treatments are available. Drinking within recommended guidelines could potentially affect the proportion of alcohol-related illness, hospitalizations and mortality seen in the population.

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Drug Misuse in the Elderly

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Substance misuse occurs mainly in young adults, with most of research focusing on this group. However, increasing age brings with it multiple pathologies and polypharmacy, with the attendant risks of dependence upon medication. There is evidence to support a growing trend to increased alcohol consumption in the over-65s, especially among women, while a generation of lifetime drug users are now entering old age¹. The need for awareness of the possibility of the existence of such a hidden disorder has never been greater in this age group. The Persian physician Avicenna described four ages of man: an age of growth, an age of prime, an age of decline and an age of decrepit old age². Those aged 65+ will include individuals in any of the latter three ages. Most research in the field has examined substance misuse by those in the declining stage.

USE, ABUSE AND HARMFUL USE

When considering drug use among the elderly it is helpful to consider substances of misuse in three broad categories: medications, both prescribed and non-prescribed; socially sanctioned psychoactive substances; and illicit substances. Religious, cultural and legal differences result in the same substances, e.g. alcohol or cannabis, being differently categorized around the world. For example, alcohol may fall within each of the three categories³. Some of the consequences of drug misuse are determined by the status of the drug rather than the physical effects of the drug. Difficulties in obtaining a drug supply and financing that use may account for as much harm as the physical effects of the drugs themselves in younger adults. Among the elderly, drugs from the medicines category are over-represented in cases of misuse when compared to other age groups. This reflects the increased access to medicines among this group, allied to the physical and social barriers that make accessing other drugs harder for this group.

At an intuitive level the clinician may balk at categorizing patients with poor compliance with misusers of non-therapeutic drugs. The aetiology, social perception and treatment approaches of the two groups differ markedly. This chapter will focus on drug misusers who display "harmful use". This is defined as:

A pattern of psychoactive drug use that causes damage to health, either mental or physical⁸.

This definition allows consideration of individuals suffering damage as a result of drug use, irrespective of the nature or the source of the drug of abuse. It excludes cases where omission of a psychoactive medication may be harmful, e.g. in cases of under-use of antidepressants.

Harmful use may be related to a single episode of drug misuse resulting in harm, such as a fall while intoxicated. More often it is a chronic condition associated with a dependence syndrome. "Dependence syndrome" describes the cluster of cognitive, behavioural and physical phenomena that are observed when use of a substance becomes a greater priority for the individual than other previously more valued activities. It is characterized by:

- A compulsion to take the substance.
- Difficulties in controlling the substance use in terms of timing and levels of use.
- Withdrawal symptoms on discontinuation of the substance, with relief of these symptoms on reinstatement of use.
- Tolerance or neuroadaptation, where increasing amounts of the substance are required to achieve effects previously possible at lower doses.
- Progressive neglect of alternative activities, due to prioritization of drug-related behaviour.
- Persistent use of the substance in spite of evidence of harmful consequences.

Presence of three or more of the above features simultaneously in the last year supports a definite diagnosis of dependence syndrome, using World Health Organization criteria⁹.

PHARMACOKINETICS

Ageing is associated with a series of physiological changes that significantly alter the fraction of an ingested drug available for a psychoactive effect. Drug absorption shows little variation with age, despite changes in gastrointestinal motility and acidity, reduced absorption surface and slowed gastric emptying. However, once absorbed, the volume of distribution in an elderly subject is likely to have changed.

Ageing results in an increase in percentage body fat and a fall in total body water. Hydrophilic drugs, such as alcohol, are distributed in body water, such that with increasing age the volume of distribution falls and the peak concentration for a given dose may rise by 20%¹⁰, resulting in lower levels of intake giving the same intoxicant effect. Conversely, lipophilic drugs, such as benzodiazepines and other psychotropics, that are stored in fatty tissue will remain in the body for longer but at lower peak concentrations. A fall in plasma albumin in old age results in increased bioavailability of protein-bound drugs, such as warfarin and diazepam.

Drug elimination occurs primarily through direct excretion or metabolism. Both routes are reduced in the elderly. Glomerular

filtration rates fall steadily in old age, leading to the accumulation of renally excreted drugs. This may be compounded by renal damage due to drug misuse, e.g. analgesic abuse¹¹. Hepatic metabolism is impaired due to a loss of liver mass and a reduced blood flow, which may also be compounded by toxic drug effects, such as alcohol leading to fatty liver. The efficiency of microsomal oxidation also falls with age, leading to reduced drug excretion of hepatically metabolized drugs¹². The combination of these effects may greatly alter pharmacokinetics in the elderly. For example, Klotz estimated the half-life of diazepam in the very elderly to be over 3 days, compared with 20 h in a younger subject¹³.

Multiple drug use increases the difficulty of prediction of the behaviour pattern of an individual substance, due to competition for binding sites and metabolic pathways. Polypharmacy may have different effects, depending on whether it is acute or chronic. Alcohol will inhibit microsomal enzyme activity in acute use, while prolonged administration will induce the same enzymes. Hence, alcohol will acutely raise concentrations of benzodiazepines, while lowering them if used chronically¹⁴.

Pharmacodynamics also alter in the elderly. Sensitivity to drugs, particularly those acting on the central nervous system, tends to increase, while drug receptor populations also change with increased age. The particular effect of the changes depends in part upon whether the receptor involved is facilitatory or inhibitory.

As a consequence of all these variables, the prediction of a drug's effects in the elderly, based on observation of its effects on younger adults, is foolhardy. Similarly rigid application of recommended "safe levels" of substance use, such as those issued by the Department of Health in the UK, may result in false reassurance to clinicians and patients with a consequent failure to identify cases of harmful use in the elderly.

CONCLUSION

The terms "old age" and "substance misuse" are both terms that have a wide range of meaning to different readers. The current literature is based primarily upon chronological age banding of individuals, as opposed to banding by overall health, possibly a more valid measure. Definitions of substance misuse are similarly varied. Often in transgenerational studies definitions of caseness are set at a level to prevent false-positive reports for younger adults. In older age groups, where less of a substance may have a greater effect, there is the possibility of missing cases if such standards are applied. The greater likelihood of drug interactions in the elderly should be considered when determining the dependency potential of any given drug or medication.

PREVALENCE AND CORRELATES

The elderly may display harmful use of any psychoactive substance. Misuse of alcohol, opioids, cannabinoids, sedatives, stimulants, hallucinogens and tobacco are all reported among the elderly. However, access to a potential substance of abuse is key to determining what an individual may misuse. Alcohol is obtainable with ease in most industrialized nations and is a socially acceptable and accessible psychoactive drug. Amongst the elderly, ill-health is common. Sedatives, hypnotics and analgesics are easily accessible through prescription and consequently, along with alcohol, are responsible for the majority of cases of harmful use. Over-the-counter medication is also easily obtained and may be misused. Illicit drugs are usually only available in potentially dangerous environments from individuals who may pose a significant risk to vulnerable older adults. Illicit drug use is not commonly observed in the elderly.

BENZODIAZEPINES

Benzodiazepines replaced barbiturates as the mainstay of pharmacological interventions in both anxiety and sleep disturbance. They maintain their relative dominance in this field despite the recent development of newer drugs with reportedly less addictive potential. Benzodiazepines accumulate more readily in the elderly due to changes in body composition, leading to a greater volume of distribution for lipophilic drugs. Chronic use may contribute to toxic effects, including cognitive impairment, poor attention and anterograde amnesia, cerebellar signs such as ataxia, dysarthria, tremor, impaired coordination and drowsiness³⁹. Increased falls and hip fractures are associated with benzodiazepine use⁴⁰, whilst withdrawal may be accompanied by rebound insomnia, agitation, convulsions and an acute confusional state. If benzodiazepines are required then short-term use of low doses of short- or medium-acting drugs is advised. There is no "safe" period of use but tolerance and dependence levels increase with prolonged use⁴¹.

Prevalence of Benzodiazepine Use

Establishing levels of benzodiazepine use is subject to the same difficulties as establishing alcohol use except where it is a prescription medication, when some idea of identity and demographic characteristics of the potential user should be available. In areas where benzodiazepines are available over the counter, the nature of users and misusers is harder to establish. National prescription audits can reflect trends in use but are unhelpful when considering particular population subgroups. Prescribing of benzodiazepines in England and Wales has fallen from 20.6 million prescriptions in 1987 to 13.9 million in 1996⁴², a fall of 32%. In England in 1996, 55% of prescriptions for benzodiazepines were issued to patients over the age of 60. Many of these prescriptions were issued to long-term users. A recent community follow-up study of 5000 over-65s in Manchester⁴³ revealed that 10% were using benzodiazepines on first assessment and that of these some 70% were taking a benzodiazepine 2 years later. A further 4 year follow-up revealed that 69% of these were still on benzodiazepines. Patients entering the study on benzodiazepines had a 52% chance of taking benzodiazepines throughout the 4 year period. Women were twice as likely to be taking a benzodiazepine as men at any stage in the study. In the USA, a study found 6.3% of a large sample of over-65s used a hypnotic, one-third of these daily and nine-tenths for at least 1 year⁴⁴. Five year follow-up found 46.6% still using hypnotics, but with a switch away from barbiturates and longer-acting benzodiazepines towards short-acting ones⁴⁵.

Use of benzodiazepines in institutional samples has traditionally been higher and associated with female gender, greater age, bereavement and poor health⁴⁶. In the USA it has been shown that one-fifth of nursing-home residents were taking potentially addictive drugs on a daily basis. The medication in question is usually a benzodiazepine⁴⁷. Studies from other countries reveal similarly high levels of benzodiazepine use among institutionalized older adults⁴⁸. The level of morbidity among institutional residents is likely to be higher than community-dwelling elders. It is unclear whether this morbidity is sufficient to explain a doubling in levels of use of benzodiazepines in this group. While chronic pain may require treatment with dependence-inducing medication, there are few indications for long-term benzodiazepine use. It has been argued that the regular use of benzodiazepines in institutions is a form of behavioural control, used more for the benefit of staff and others than these users. In many cases, the individual may be incapable of giving valid consent to taking such medication. The use of medication in such circumstances may be

considered benzodiazepine misuse by some and as elder abuse by others⁴⁹.

Correlates

Psychiatric Morbidity

Significantly high rates of psychiatric disorder have been described among elderly benzodiazepine users⁵⁰. Among elders using short-acting benzodiazepines as hypnotics, one-third reach caseness for depression, while a further one-third have a diagnosable anxiety disorder. Amongst users of anxiolytic benzodiazepines, half are depressed and one-fifth are anxious in spite of treatment. These results are not evidence of a causative relationship, although the most likely indications for initiation of such medication by a prescriber are likely to be presentation with such symptoms. As with alcohol misusers, one-third of elders requiring inpatient treatment for benzodiazepine misuse are of late onset, while two-thirds have graduated from misusing benzodiazepines or other drugs whilst younger⁵¹. The incidence of co-morbid alcohol abuse has not been consistently shown to be significantly greater among benzodiazepine misusers^{51,52}. An all-age study found that DSM-III-R Axis I co-morbidity existed in all cases of a sample of benzodiazepine dependent users in Spain⁵³. The commonest diagnoses were insomnia, anxiety disorders and affective disorders. Obsessive-compulsive, histrionic and dependent personality disorders were found in half the cases and physical problems in one-third of cases.

Gender and Age

Benzodiazepine use is over-represented among women of all ages. The likelihood of use of a benzodiazepine increases with age. There is little evidence that this gender divide narrows on reaching old age. Legislative approaches and prescribing guidelines have made some inroads into the over-representation of prescribing to the elderly⁵³. Increasing public awareness of the side effects of benzodiazepines and an increase in advocacy services for the elderly are likely to have a similar effect.

Other Prescribed and Over-the-counter Medication

As indicated earlier, the elderly routinely receive a wide variety of medications, the majority of which may be misused. A quarter to half of the elderly experience chronic pain. In acute use the dependency potential of analgesics is believed to be around 0.1%. In chronic conditions the situation is somewhat different. Ten per cent of over-64s are on prescribed analgesics at any one time, with at least an equal number using over-the-counter medication. Edwards and Salib⁵⁴ found 3% of a community sample of over-65s to have been using mild opiate analgesics for a period of at least 1 year; 40% of this group were deemed to fulfil the criteria for dependence upon these drugs, with dependence levels as high as two-thirds among users of co-proxamol.

In addition to the dependence caused by these drugs, physical harm may also result, e.g. nephropathy may be caused by the use of paracetamol, salicylates and pyrazole derivatives, while renal impairment occurs with non-steroidal anti-inflammatory drug use⁵⁵. Chronic nephropathy may also be caused by the excessive ingestion of analgesic mixtures combining two or more antipyretic analgesics, along with codeine or caffeine (both independently capable of causing addiction). Such acute and chronic effects are more likely amongst the elderly, where relative drug levels are higher and less biological reserve exists. Similar physical

complications may arise from the misuse of other medications, the commonest being laxatives and cough mixtures.

ILLICIT DRUG MISUSE

Little is known about levels of illicit drug use among the over-65s, although the general perception is that it has been less of a problem than the misuse of prescribed medication. In the Epidemiological Catchment Area Study (ECA), only 0.1% of elders met the criteria for drug abuse for an illicit substance in the previous month⁵⁶. Lifetime prevalence was 1.6% for over-65s. This may change as younger generations with a pattern of recreational drug use reach old age. In the UK, few cases of illicit drug use among the over-65s have found their way into the literature; one exception is a series of seven elderly reported to have initiated injecting heroin in later life. They attributed their behaviour to a combination of loneliness and depression⁵⁷. In the USA, in a recent study of a Veterans' Administration old age psychiatry inpatient facility, 3% of the patients were found to have a primary drug misuse disorder involving prescribed medication, while 1% were addicted to illicit substances⁵⁸. Also in the USA, attendance at methadone maintenance clinics by the elderly is reported to be rising, although over-60s still form 2% of those attending⁵⁹. Similarly, a number of elders are reported to continue their use of cannabis into late life⁴⁷. Anecdotal evidence also points to some individuals initiating the use of cannabis in later life in a search for its reputed therapeutic benefit in conditions such as disseminated sclerosis.

On balance, it appears that illicit drug use is less of a problem in the elderly than the abuse of legally sanctioned drugs. It remains to be seen whether individuals currently abusing illicit substances in younger age groups carry this behaviour over into old age. The nature of the subject has not lent itself to prospective studies as yet. One might expect greater levels of illicit drug use in future generations of older adults, although difficulties associated with obtaining a supply of such drugs with increasing infirmity are likely to account for some cessation in use. It is also tempting to speculate that those abusing illicit drugs as younger adults may switch to misusing prescribed medication in later life.

POLYSUBSTANCE MISUSE

The elderly have access to a variety of drugs of misuse. In many cases they may misuse one drug without misusing others. This is often the case with prescribed medication, where one medication is overused while compliance with the prescription is maintained for the others. Where non-prescribed substances become involved, the possibility of abuse of more than one substance is elevated. Finlayson⁵⁰ found 15% of over-65s requiring inpatient detoxification from alcohol were also dependent upon a second substance, usually a hypnotic, anxiolytic or analgesic.

The phenomenon of cross-tolerance must also be considered. Psychoactive substances may have a cumulative effect, due to either a shared outcome effect or to different drugs acting as interchangeable substitutes for one another (cross-tolerance). Cross-tolerance exists within each class of drug, such that the clinician should always consider the total benzodiazepine, barbiturate or opioid dose, using class-specific equivalence charts⁶⁰. Cross-tolerance for some drugs may also occur outside of the class, most notably for alcohol, chlormethiazole and benzodiazepines. While this phenomenon is widely exploited for detoxification, failure to consider the possibility of its existence may lead to overlooking cases of dependence.

Conclusion

The abuse of alcohol and of prescribed medication remain the most prevalent substance misuse problems. There is little clear evidence of great changes in individuals' addictive behaviour patterns with increasing age. When one considers the prevalence of substance misuse among younger cohorts, it would appear that, as the population ages, not only will the absolute numbers of elderly with a substance misuse problem increase but the proportion of the elderly population with such a problem will increase too.

TREATMENT

Treatment of substance misuse is a multistage process involving the integrated use of physical, psychological and social interventions. These interventions should, where possible, run concurrently as opposed to consecutively and must be provided in a form that is acceptable to the individual and sensitive to the specific needs of the elderly. Amongst this client group, individuals rarely present complaining directly of a substance misuse disorder but may present with associated physical problems, or a problem may be detected during routine consultation with health professionals or carers.

The first step of treatment is the identification of cases. This requires clinical observation allied to sensitive yet persistent enquiry. The routine use of standardized screening tools may help to focus clinical impression more accurately. Once identified as potential candidates for treatment, the patient's attitude towards his/her substance misuse requires examination. Exploration of the risks and a discussion of potential avenues for change may help to establish or reinforce the motivation to change. Drugs that cause significant physical dependence may necessitate detoxification regimens, while co-morbid conditions such as depression that perpetuate the disorder need to be adequately treated. Social issues, such as housing and a social network that comprises mainly of substance misusers, may perpetuate the problem and need to be examined for opportunities to change. The individual requires psychological rehabilitation to address the issues that may have contributed to the uncontrolled use of substances and to provide future coping strategies to prevent a relapse into substance misuse. While these approaches apply to all substances of misuse, the majority of research has focused on alcohol misuse.

DETECTION

Self-presentation by elders may be limited by a number of factors⁶¹. Practical issues, such as accessibility of treatment centres to disabled individuals, large print information sources for the visually impaired and the availability of domiciliary treatment, are fundamental. Elders may not realize that they are ill, or may not realize that the medical profession identifies substance misuse as an illness and will offer help. Traditional forms of service promotion may fail to reach the elderly, while a service staffed by young professionals may seem intimidating or inappropriate for someone much older, particularly if his/her substance misuse is associated with a high degree of shame as is often the case in this group.

If self-presentation is unlikely, then the number of professional caregiver contacts that the elderly have provides a further opportunity for education of individuals about the problem and potential sources of help. This resource appears underdeveloped at present, with a need for better training for carers in identification of at-risk individuals and in appropriate actions once misusers are identified⁶². Currently, evidence suggests that

carers are often unaware of sources of help and frequently are in collusion with alcohol misuse, citing reasons such as the elder "has not got long to live" or that "it's his only pleasure"⁶³. Studies have found that many agencies providing care for the elderly have no written policy to guide their employees when encountering a client with an alcohol problem. Greenwood⁶⁴ argues that substance misusers, and the elderly in particular, suffer as the result of stigmatization, as their disorder is perceived as self-inflicted and with a poor outcome prognostically. This stigmatization may contribute to difficulties in communication and empathy on the part of some caregivers, including doctors. This stigma may be reflected in a clinician's reluctance to become involved by acknowledging a problem. For other carers, their own previous experiences with elderly substance misusers, both professional and personal, may lead to attempts to justify the behaviour, resulting in a similar loss of objectivity.

BENZODIAZEPINE USE DETECTION

Prevention and early recognition form the basis of management of benzodiazepine and other drug misuse among the elderly. Appropriate prescribing of sedatives for time-limited periods should be accompanied with vigilance for drug-seeking behaviours. Such behaviours include early requests for repeat prescriptions or requests for increased doses and should be regarded with suspicion. The elderly may also receive medication from multiple sources, particularly where they are under the care of prescribing hospital specialists as well as their primary prescribers. Careful exchange of clinical information is vital in such settings. For those abusing over-the-counter medication, chance presentation or the intervention of a pharmacist present the best hope of detection.

No screening tools have been validated to detect cases of elder benzodiazepine use. While urine screening provides a reliable means of establishing the presence or absence of drug metabolites, its clinical utility is limited by the unacceptability of the test to many who may be offended by the suggestion that they have a substance misuse problem. Second, urine screening is usually qualitative rather than quantitative. For those abusing a prescribed drug, the mere presence or absence of the drug is clinically uninformative. Individual variation in pharmacokinetics makes qualitative testing unreliable. With these considerations in mind, the need for dependence-inducing drug prescriptions should be regularly reviewed and co-morbid contributory conditions, such as depression, should be actively treated. Changes in legislation on prescribing practice may reduce the opportunity for drug misuse⁷¹.

INITIATING TREATMENT

There are no published data about the level of uptake of offers of help once elders abusing substances are identified. Motivational interviewing and education as to the risks of alcohol use, along with the benefits of even a small reduction in levels of alcohol intake, may persuade some elders to change. Unfortunately, the pessimistic attitudes held by many professionals and carers towards the likelihood of successful resolution of the problem are frequently also held by the individual too. A fatalistic resignation to a life of substance misuse is often reported, particularly by long-term users, while more recent-onset users may express greater motivation for treatment⁷².

Once long-term use of benzodiazepines is established, dose reduction can be difficult to achieve. Withdrawal insomnia and rebound anxiety make patient motivation difficult to achieve. Where abstinence is desired, a conversion to a medium-acting

benzodiazepine and a gradual reduction in dosage over the course of months is advisable. Rapid detoxification is associated with breakthrough withdrawal symptoms and may be complicated by convulsions. If a rapid withdrawal is necessary, it is best conducted in an inpatient setting if severe dependency is suspected. As with alcohol, the withdrawal period for the elderly is more likely to be complicated by confusion than in younger adults.

In cases where abstinence is not achievable or desirable, a minimization of dose and adoption of a non-daily pattern of use are reasonable targets. Psychological techniques, such as relaxation training and educative initiatives in the areas of sleep hygiene and correct medication use, may also prove valuable. Cormack⁸¹ demonstrated that writing to benzodiazepine users in primary care urging them to reduce their medication use resulted in a fall in total use by one-third over the next 6 months.

Treatment of other forms of drug misuse in the elderly is under-researched. Anticipation of problems and safe prescribing remain paramount in treatment and prevention. Misuse of analgesics may require formal detoxification if opioids are involved or physical dependence has developed. More often the patient requires information to allow him/her to make an informed choice about drug use and an alternative form of treatment for his/her condition. Still less information is available on the treatment of illicit drug use in the elderly. At present there is no evidence to suggest that an approach other than that used for younger adults should be adopted, although adaptations of such treatments should involve lower doses of medication and the adoption of a less directly confrontational approach.

PSYCHOLOGICAL INTERVENTIONS

Once a patient is detoxified, rehabilitation is necessary to address the issues behind his/her substance use and to foster coping strategies for the future. Few studies have examined the particular needs of the elderly in a rehabilitation setting. Janik and Dunham report on comparative outcomes for over 3000 over-60-year-olds and younger entrants into alcohol treatment programmes⁸². Outcome measured in terms of alcohol intake, therapist assessment and alcohol problems after 6 months showed no differences between the groups.

Outcomes from programmes designed specifically with the elderly in mind may be more appropriate for consideration. Some success has been claimed for models encouraging the development of social networks with self-management skills⁸³. Kofod⁸⁴, in a small study, reported that retention in outpatient treatment of older adults was greater in an age-specific treatment group that focused on socialization and minimal confrontation (a mainstay of many programmes), compared with older patients in a mixed-age treatment group. At 1 year follow-up the effect was lost. Variations of the Alcoholics Anonymous 12-step model tailored to the needs of elders have been reported upon⁸⁵ in the USA, with varying degrees of success. Models low on confrontation, traditionally regarded as fundamental to overcoming denial on the part of the patient, appear to be supported by the work of Kashner⁸⁶, who found that 1 year follow-up of elders in a confrontational programme revealed half the levels of abstinence as compared with a group in a programme where self-esteem, tolerance and peer relationships were promoted. Behavioural approaches, including cue identification and avoidance, have also been reported to be of clinical benefit⁸³. A programme focusing on cognitive techniques, such as cognitive restructuring, assertion training and self-monitoring of drinking, resulted in 75% of those completing the programme sticking to their treatment goals at 1 year follow-up. The evidence suggests that a range of therapeutic techniques may be beneficial for the elderly and that local

provision may depend upon the skills available to the treating agency. It is suggested that even if an elder's only therapeutic programme is not available, a better therapeutic outcome may occur from a more homogeneous group, where the opportunity for identification and vicarious learning is enhanced.

The above studies all relate to the outcome of alcohol treatment programmes. Even less age-specific studies are available to guide the clinician in the provision of aftercare to the elderly non-alcoholic drug user. An avoidance of drugs that have a dependence potential is advisable if practical. Adequate rehabilitation and continuing support of the individual are indicated. This may be provided through generic old age psychiatry services or through specialist drug services, depending upon which service appears best able to cater for the specific needs of the user. The choice of service provider should reflect the lifestyle of the patient, as opposed to being a decision based solely on chronological age. Further services may also be available in the form of mutual support groups similar to those available for alcohol. The adoption of a cognitively-based programme low on confrontation and designed to foster strong social support appears optimal, as shown in work in the field of alcohol.

Conclusion

Substance misuse and old age psychiatry have long been unpopular choices for specialization. Both fields are known for providing challenging patients with differing priorities to those of the clinician. Research in either field is hampered by the difficulty in obtaining reliable clinical data on conditions for which few empirical measures exist. The field of old age substance misuse has suffered to some extent in clinical practice, where patients do not fit neatly into either service and welcomed by neither.

It is, however, clear that there exists a significant morbidity due to drug use in the elderly. The problem may be iatrogenic and autogenic in origin. Increased life expectancy and the cohort effect of generations of recreational drug users reaching old age are likely to intensify the problem. Adequate research to identify at-risk individuals and the provision of appropriate and accessible treatment services for the elderly drug misuser remain one of the major challenges to health care providers at the start of the new millennium.

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Benzodiazepine Use and Abuse in the Community: Liverpool Studies

Kenneth C. M. Wilson and Pat Mottram

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Benzodiazepine use in older people is associated with increased falls¹ and psychomotor changes, leading to the recommendation that they should rarely be prescribed². Despite the fact that they may play an important role in the treatment of some psychiatric conditions³, it is evident that they should be used with caution in this age group.

Recent epidemiological studies in Liverpool clearly demonstrate cause for concern. Between 1982 and 1983 we examined the prevalence of mental illness and the use of drugs in a random sample of 1070 community residents aged 65+⁴. A similar study was conducted on a larger age- and gender-stratified sample ($n = 5222$) of the same age group between the years 1989 and 1991⁵. The Geriatric Mental State Examination was used in both studies. The instrument also collects data concerning prescribed and non-prescribed drug use over the month preceding the interview.

The results of the two studies have been compared in some detail⁶. The overall prevalence of benzodiazepine use was 12.8 (CI: 10.8–14.8) in 1982–1983 and 10.8 (CI: 9.9–11.6) 10 years later, indicating no significant change in drug use prevalence. Over the 3 years of the later study, 2.5% of non-users started taking benzodiazepines. Over two-thirds of all users were still taking benzodiazepines 3 years later. Analyses by mental state demonstrated that the largest proportion of people receiving benzodiazepines were depressed. People aged 80+ were three times more likely to be users than those aged 65–70. Both depression and anxiety were risk factors. Subsequent analyses of the data⁷ examined the use of benzodiazepines in the context of depression. It is evident that nearly twice as many depressed older

people were users of benzodiazepines than antidepressants. This may be explained by the increased emphasis on agitation, initial insomnia⁸ and the relative under-reporting of depressed mood in older depressed people⁹.

The findings of these studies should be viewed with some caution in view of the relatively long period between interviews in each study. It is self-evident that users may be intermittent in their use of benzodiazepines and other medications. However, the larger study incorporated questions concerning compliance. Data were available from 203/208 benzodiazepine users at their last interview. Of these, 89.6% stated that they were taking drugs as prescribed, 4.4% occasionally missed a dose and 5.4% often missed a dose. Only one stated that he/she had stopped taking the prescribed medication.

The findings of these studies imply that approximately 10% of older community residents are prescribed and taking benzodiazepines. Benzodiazepines are frequently prescribed to the very old and depressed, the majority of whom remain depressed for a significant period of time. Once commenced on benzodiazepines the majority take them for at least 3 years. There does not appear to be any change in prescribing habits across the interim period of 10 years.

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Part K

Learning and Behavioural Disorders

Old Age and Learning Disability

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The age structure of people with learning disabilities is changing and more are surviving into old age, indicating considerable achievements in health and social development.

Historically, there was little provision for the needs of older persons with learning disability. The reason for this was that such persons had a short lifespan and many died before reaching old age, for reasons such as inadequate medical treatment and the life-threatening complications associated with their condition. During recent years, the provision of formal care for adults with learning disabilities has changed from a largely institutional service to an increasingly community-based one. This has affected the collection of health statistics. Hospital populations provided the main source of data in the past. Although registers of need are a statutory requirement for children in the UK, there is no similar requirement for adults with disabilities. However, many local authorities do maintain learning disabilities registers, and include a wider range of service users than those who would previously have been long-stay hospital patients.

SOCIAL BACKGROUND

In the mid- to late 1800s, "mentally defectives" (including people with learning disabilities and those with mental illness) were kept away from society and were housed in asylums. One reason for the institutionalization of mildly learning-disabled people can be found in the social turmoil of the early 1900s. Young adults were placed in institutions to curb their sexuality, as it was thought that their offspring would have intellectual limitations and would themselves have low-IQ children, and that eventually the national IQ would fall. After the menopause, many of the women were allowed out on licence, working as ladies' maids, for example, and returning to the asylum when the problem of old age meant they were of no further use. Many were of normal intelligence, brought into institutions because of adverse social circumstances and detained there. After World War II, life expectancy began to improve within the institutions with better medical care and more positive lifestyles^{2,3}. Mildly disabled older adults were highly valued as a substantial work force in the institution.

In 1959, a review of the Mental Health Act allowed patients an informal status and many became voluntary patients and discharged themselves, made their way into the world and were lost to helping agencies and statistics. Others stayed on in institutions, probably because of infirmity or institutionalization. Many of these will have been discharged as elderly individuals in the wave of community care in the 1980s, promoted by the philosophy of normalization and governmental encouragement. Wolf and Wright³ found that younger, more able people were

more readily discharged into the community and the older and more disabled population remained in institutions.

CAUSES OF DEATH

People with learning disabilities are living longer overall but the level of disability is correlated with longevity and this association is more marked in the earlier years. Improved standards of care, higher expectations and more positive attitudes to treatment for serious illness have contributed to increasing longevity. Prolonged survival is now the norm, even for the most severely disabled children. In people with Down's syndrome, for example, heart disease is now treated surgically if indicated and overall medical care is much improved. The overall increased longevity means that disorders such as dementia and cancer have become more prevalent. There is a tendency for profoundly learning-disabled people to be more seriously physically disabled and to have more health problems. Even with better care, the life expectancy of profoundly and multiply disabled people is still reduced. The mortality rate for people with learning disabilities is higher compared with the general population. Carter and Jancar^{4,5} studied a hospital population and found marked changes in the causes of death over a 50 year period. Prior to 1955, tuberculosis was a major cause of death, and after 1955 non-tubercular respiratory infection accounted for 46% of all deaths examined. Other identified causes were myocardial infarction, cerebrovascular accidents, pulmonary embolism and status epilepticus, which accounted for 15%. Epilepsy is both more prevalent and more likely to occur in a poorly controlled form with significantly increased mortality². Hollins *et al.*⁶ reported that death certificates were not a reliable source of data about cause of death and that learning disabilities were rarely mentioned, with respiratory disease being the major cause of death, suggesting a failure to recognize underlying medical conditions. Those with mild disabilities have lifespans close to those of the general population, dying from cardiovascular, neoplastic and terminal infections, similar to those of elderly people of originally normal intelligence⁷.

FREQUENCY OF DEMENTIA IN PEOPLE WITH LEARNING DISABILITY

Epidemiological studies, whether cross-sectional or longitudinal, are often difficult to do. The assessment of the premorbid state is affected by limited educational opportunities, social deprivation and the low expectations of many people with learning disabilities. The true intellectual functioning of the person with a learning

disability at baseline is often difficult to measure¹. Studies suggest that dementia is about as common in the learning-disabled population as in the general population and has the same range of clinical phenomena, provided that each patient's unique intellectual baseline is allowed for⁸. Cooper⁹ surveyed a population of elderly people with learning disabilities and found that 22% had dementia; however, the small number precluded analysis of subtypes. An earlier survey by Reid *et al.*¹⁵ found a prevalence of 13.6%. Tait¹⁰ found a prevalence of dementia similar to that in the general population.

THE MEDICAL CONTRIBUTION TO THE DIAGNOSIS OF DEMENTIA

Dementia can only be diagnosed after a careful history is taken and an examination made, the premorbid and presenting personalities assessed and other reasons for deterioration excluded. Long-standing visual and hearing problems are common in people with learning disabilities and may only be discovered as sensory deficits increase and further affect functioning¹¹. Hypothyroidism, deterioration caused by inappropriate medication, communication disorders and psychiatric illness, both organic and functional, may all cause pseudodementia. Depression is often precipitated by loss of caregivers or familiar environments and may present, in addition, as behavioural and personality change. Symptoms such as incontinence may be related to the inappropriate architecture of the residence or to shortage of staff.

A detailed clinical examination will highlight dental and chiropody needs, general medical disorders and complications of long-standing disability. Consideration can be given to the need for aids and appliances to minimize the deficits and ease the burden for carers. People with learning disabilities rarely have access to health education or health promotion, so that screening for anaemia, hypertension, glaucoma or carcinoma of breast or cervix will hardly ever have been done.

The examination of someone with learning disabilities who may have superimposed dementia may therefore offer the opportunity to put right some of the deficiencies of older-style services, identify current social dilemmas, diagnose dementia in the context of long-standing deficits and consider, with the caregivers and the multidisciplinary team, how needs can be met¹².

Thorough assessment will provide information on the following:

1. The developmental intellectual disability.
2. Other long-standing disabilities and comorbid conditions.
3. Long-standing psychiatric illness and behaviour disorder.
4. Illness and disability superimposed and due to ageing.
5. Psychiatric illness associated with old age.
6. Dementia, if present.
7. The skills and needs of the person.
8. Recent life experiences.
9. The patient's wishes and those of the caregiver for residential and social care.

DEMENTIA AND DOWN'S SYNDROME

The association between ageing in adults with Down's syndrome and the development of dementia attracts interest from both researchers and clinicians. There is now a substantial literature on the genetic link between Down's syndrome and Alzheimer's disease. Several studies, using a variety of diagnostic criteria, have reported increasing age-specific prevalence rates for Alzheimer's disease in people with Down's syndrome. However, whilst the

prevalence rates vary across studies, in no study has the rate reached 100%, which might be expected given the neuropathological data¹³. Dementia is often accompanied by epilepsy, loss of skills (which perhaps were not well developed in the first place) and transient behaviour problems, together with personality changes.

Families who have cared for their relative with Down's syndrome to the point where they develop dementia need support, information and the chance to talk to someone who understands the natural history of Alzheimer's disease. They may feel guilty and confused. An understanding of their confusion is required, and support is needed as they accept the diagnosis of dementia and its inevitable outcome. They may need to consider changes to their lifestyle and to look to the wider network for longer-term care and support.

WHAT OF SERVICES?

One of the most challenging ways to think about service development is through the proper consideration of the philosophy of normalization, which states that services that are highly valued and normative should be used by those who are at risk of being devalued. The difficulty is that both "the elderly" and "people with learning disabilities" are potentially devalued groups; both services tend to be underfunded and considered to be bottomless pits of needs.

The major debate is whether to use the services the rest of the population use, i.e. the geriatric or psychogeriatric services, to continue with learning disability services (improved as necessary), or to develop something new. Probably the best solution is to consider services for each individual, facilitating access to what is available and campaigning for what is not.

In England, national policy requires local authorities to offer person-centred planning to all people with learning disabilities. People living at home with elderly family carers are a priority group for receiving services and supports based on what is important for them as individuals and for receiving a regularly updated health action plan¹⁶.

REHABILITATION INTO THE COMMUNITY

The closure of long-stay mental handicap hospitals in the UK has created new social and medical dilemmas¹⁰. Some people have been in hospital for many years, and moving into the community presents many difficulties, with geriatric needs superseding developmental disabilities. Some hospitals have remained to accommodate this small group, but many hospitals have closed completely and people moved to community residential provision. Many elderly people with learning disabilities have their only network of friends on the campus of the hospital, and it is important to try and maintain this. They are usually out of touch with relatives and a return to the county or borough of origin is not always meaningful or in their best interest. With careful introduction, these people may usually be accommodated in community services and may be far more competent than typical geriatric patients¹⁴. Because their life experience is so different, it may be better to care for them in an establishment able to take several residents from the same hospital.

CARE IN THE FAMILY HOME

The elderly person with learning disability may be cared for by an ageing parent with a similar dependency level. The fragile world of a person with learning disability has often been prematurely

closed because of the declining abilities of the caregiver. With the death of remaining carers, the experience of multiple loss may precipitate depression as part of the bereavement response¹⁷. A small group home may then provide a substitute for the family home. With foresight and planning, some people may stay in their own home, with up to 24 hours support if needed.

THE PERSON WITH MULTIPLE DISABILITIES

Learning disability is known to be associated with an increased prevalence of coexisting diagnoses that also affect life expectancy, with the result that those with multiple disabilities and superimposed problems of old age are few in number. There are challenges in management which may require a combination of learning disability, psychiatric and geriatric expertise. Some hospitals have developed special units to accommodate this small group, but new services are community-based and provide a multispecialty and multidisciplinary format. Private and voluntary agencies have often led the way.

DAY CARE

Accommodation and care varying from minimal to total are only part of the needs to be met. Daily activities and social contacts will need to be provided, and this may be difficult if the person has relocated in an area where he/she is not known. Because of shortage of places in social education centres (the core provision for adults with learning disability in the community), adults with learning disabilities usually retire by 65 years and then have only the occasional part-time arrangement in day centres for the elderly, clubs, adult education institutes and religious activities. As the number of elderly people with learning disabilities in the community increases, person-centred approaches to planning individual arrangements will be more important.

COMMUNITY TEAMS

Most areas in the UK have multidisciplinary community teams for people with learning disabilities and the trend is for these specialist staff to facilitate access to assessment and management within mainstream geriatric and psychogeriatric services. The opportunities of working together are great, thereby ensuring that dividing lines in health care do not detract from whole-person medicine.

TERMINAL CARE

Elderly people with learning disabilities may die at home, in oncology or general medical wards or in a hospice. The combined community services may be needed and include the terminal care team. Chaplains, moving from long-stay hospitals into ecumenical teams in the community, may be called to minister, together with community nurses and community team members, to ensure a dignified and good death.

CONCLUSION

The majority of elderly people with learning disability now live in the community and are more likely to outlive their parents and share the experience of becoming old with the rest of the population. This is an opportunity to meet their several needs successfully and not to repeat the errors of segregation that have littered learning disability services throughout the industrial world. There is need for more research at a clinical and planning level to underpin creative services, and up-to-date knowledge available for those who do the caring and for those who help the carers.

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Elderly Offenders

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EXTENT AND PATTERN OF CRIMINAL BEHAVIOUR IN THE ELDERLY

Criminal behaviour is most common during youth and declines sharply with increasing age. Older people are responsible for only a small part of the total amount of recorded crime and it is likely that they also make a limited contribution to the “dark figure” of unrecorded offences. The lower proportion of males in the elderly population, their retired status and the absence of the risk-taking attitudes of youth may be of relevance. A review of the criminal statistics for England and Wales indicates that the elderly’s share of most offence categories is less than 1% of the total for all ages. Shoplifting is the most common indictable offence to involve the elderly and this is the only offence category where older women make a significant contribution¹. The elderly’s share of sex offences (other than rape) is higher than for other indictable offences, varying between 5% and 9% of each category in England and Wales. Schichor² notes a similar 5% figure for the elderly’s contribution to overall sex offences in the USA. Offences involving children are most common. In Craissati and McClurg’s study of a consecutive series of 356 men from two London boroughs convicted of sex offences or released on parole (1993–2001), 22 men (6.1%) were aged 60 or over, 19 men were convicted of offences against children, two against adults and one of possession of prohibited material involving children^{3,4}. The elderly contribute very little to the total number of those convicted of burglary, robbery and drug offences.

Violent behaviour in the elderly leading to conviction is extremely rare. Essex Police District recorded reports of violent incidents in only three people aged 65+ during a 14-month period⁵. In 1998–1999, 3.9% of homicides (19 cases) in England and Wales involved those aged 60+; six of these were convicted of Section 2 manslaughter (diminished responsibility), comprising 31.6% of the elderly group. In comparison, 6.8% of homicide cases where the perpetrators were under 60 resulted in a Section 2 manslaughter verdict, supporting the more prominent role of psychiatric disorder in the older offenders¹.

Kratcoski and Walker⁶ studied 82 cases of homicide committed by those aged 60+ in the USA and concluded that elderly homicides were more likely to involve a spouse or other relative as victim than the non-elderly; 81% of the offenders were male, and 15% of the elderly offenders committed suicide, the victim being the wife in each instance.

Hucker and Ben Aron⁷ compared 16 elderly violent offenders with a group of young violent offenders and with another group of non-violent elderly sex offenders, all selected by referral to a psychiatric clinic in a case note study; 69% of the elderly violent group were diagnosed as having either an organic brain syndrome or a functional psychosis, compared with only 19% of the non-

violent elderly and none of the younger group. Paranoid symptoms were prominent in the elderly violent group, irrespective of diagnosis.

MENTAL ABNORMALITY AND OFFENDING IN OLD AGE

The most common associations of criminal behaviour in the elderly are alcohol abuse⁸, homelessness and psychiatric illness⁹. In an early paper on this topic, Norwood East¹⁰ suggested, on the basis of his clinical experience in prisons, that the possibility of psychiatric disorder should always be considered in those who offended for the first time in old age. Particular attention has also been drawn to first-offender shoplifters¹¹, although the contribution this disturbed group make to the whole is unknown.

In a study of 153 referrals aged 60+ to the community services branch of Essex police¹², 97 had been apprehended for shoplifting. The prevalence of psychiatric disorder in the 50 people interviewed was higher than in other community samples; 38% of those charged with shoplifting were identified as cases, with 9/11 cases belonging to AGE-CAT organic and depressive syndrome groups¹³. In a study of men remanded in custody, Taylor and Parrott⁹ found that nearly 3% of men were aged 55+. Half of these men had active symptoms of psychiatric disorder and half some form of physical disorder—twice the rates of those under 55. The commonest psychiatric problems were alcoholism (27%) and major functional psychosis (37%). Less than a quarter of the over-65s had a permanent address, suggesting that homelessness was an important determinant of custodial remand. Twenty-four of 1062 restricted patients admitted to hospital in 1998 in England and Wales were aged 60+; nine of these patients had been convicted of homicide, six of other violent offences and two of sexual offences¹⁴.

CLINICAL ASPECTS OF OFFENDING IN THE ELDERLY

Affective Disorder

Roth¹⁵ noted that the rare violent acts committed by aged men often arise in a setting of depressive illness with suicidal ideation. Depressive homicide most often involves a man killing his wife. The killing is generally viewed by the patient as at least partly altruistic, on account of thoughts that she would be unable to cope without him, although a dynamic formulation may indicate unacknowledged hostility. It is common for the act to be followed

by suicide. Knight described several cases, noting that there is generally no history of previous violence or discord¹⁶.

The disinhibition and irritability of mania may also lead to aggressive behaviour, although serious violence is uncommon. Both types of mood disorder may be associated with shoplifting, on account of poor concentration, diminished concern about social rules or associated dynamic factors. In relation to shoplifting, an emotional disturbance following a life crisis but falling short of depressive illness may form the background to the offence. It should also be borne in mind that the trauma of arrest itself in an elderly person with no previous symptoms may lead to psychiatric sequelae, including suicidal ideation.

Case History 1: Depressive Homicide

A 68-year-old man with previous episodes of depressive illness developed depressed mood and agitation over several weeks. There was no history of violence and he was known as a quiet kindly man. He became preoccupied with two themes, first that the water supply in his house was infected and had caused his wife's psoriasis, and second with guilt regarding a minor sexual misdemeanour that had occurred during his teens. His wife asked him to just try and do the shopping, something he found particularly difficult when depressed. He experienced the sudden thought that if he killed her, the action would have the dual benefit of releasing his wife from her misery and ensuring that he received a life sentence, a fitting punishment for the sexual misdemeanour. He strangled her and rang the police. He was bailed to hospital and later placed on probation, with a condition of psychiatric treatment.

Case History 2: Shoplifting and Depression

A 75-year-old retired plasterer with no previous convictions received the news that his son-in-law was having an affair. He himself had endured a stressful marriage for a lifetime. He became depressed in mood and overwhelmed by intrusive, angry thoughts towards his son-in-law. His sleep became disturbed and his powers of concentration diminished. While shopping he was observed to pick up a paint brush and slip it into his jacket. He then clambered over a barrier to leave the store. The shop refused to stop proceedings, but the Court dismissed the charge on hearing psychiatric evidence. The emotional crisis subsided with time and a period of supportive counselling.

Schizophrenia

The most common offence associated with schizophrenia in those that are homeless is shoplifting. However, both violent offences and arson may occur, on account of paranoid ideation.

Case History 3: Violence Associated with Paranoid Delusions

A 75-year-old man was arrested following his approaching a group in his local pub, grabbing one person by the neck and cutting his lower throat with a carving knife. The publican reported that the elderly man was a moderate drinker and that he had been expressing abnormal ideas over the previous year. He had had a partial pneumonectomy for lung cancer 3 years prior to the offence. At interview he said that neighbours watched him all the time, made derogatory remarks about him and had a laser machine which they used on his body. On the day of the offence, he said he had heard one of the group of men say "He stinks", and

concluded that they were associated with those acting against him. He returned home for a knife and said later he only wished to frighten the victim. No evidence of recurrence of his lung cancer was found at that stage. Following treatment with antipsychotic medication, he no longer complained of abnormal experiences, although he retained some delusional ideas. He became relaxed and content in contrast to his agitated state prior to treatment and said that he had no desire to revenge himself on the people involved in the paranoid ideation. He pleaded guilty to wounding with intent and was placed on a hospital order under Section 37 of the Mental Health Act (1983), with restrictions on discharge under Section 41. There were no management problems, although he remained in hospital until his death with metastases 18 months later.

Dementia

Assaultive behaviour in dementia is not uncommon, although within a family or institutional setting the behaviour is unlikely to form the basis of a formal charge. Disinhibition, misinterpretation and the pressures of close living with others may be contributory factors. Petrie *et al.*¹⁷ studied a series of 222 consecutive admissions to a psychogeriatric unit in the USA and noted that 139 had shown verbal aggression or violence, 18 incidents involving the use of knives or guns; 39 of these patients were suffering from senile dementia and the remainder from functional psychoses. Disinhibited sexual behaviour or fire-setting may also reflect the loss of cortical inhibiting factors in dementia.

Persistent theft from shops may be linked with absent-mindedness in the early stages of illness or with a more general deterioration in social behaviour at a later stage. Mendez described a case of persistent stealing in a 71-year-old man with dementia who constantly picked up small items with no explanation for his actions¹⁸.

Alcoholism

In addition to alcohol-related offences, such as driving while intoxicated, problem drinking may be associated with theft, criminal damage, violent or sex offences. Many cases involve a variety of factors, e.g. about one-third of elderly homicide offenders in a series studying coroners' files had been drinking at the time of the killing⁶.

ASSESSMENT OF THE ELDERLY OFFENDER

A careful assessment of mental and physical health and of the person's social circumstances and the quality of relationships within this is necessary. Particular note should be taken of the use of alcohol and of prescribed or proprietary medication. Where the charge is more serious, full details of the allegation should be obtained from witness statements and informants.

In assessing a defendant's fitness to plead, consideration is particularly given to the defendant's ability to understand the nature of the charge and the significance of his plea and to follow the process of the trial. In some cases, discontinuation of proceedings may be more appropriate.

THE ELDERLY WITHIN THE CRIMINAL JUSTICE SYSTEM

Appropriate assessment that would identify those offenders requiring social and psychiatric intervention is often lacking,

although there has been a greater emphasis on diversion from the criminal justice system in recent years. Cautioning is as common a disposal as conviction in England and Wales for all but the most serious crimes in those aged 60 +⁵. In addition, many shops have adopted policies of not reporting shoplifting in those of retirement age, preferring to ban persistent offenders from their premises. In the USA a number of schemes have been developed to deal with the older offender, both pre-trial and after sentencing¹⁷.

In the Essex cohort of 153 elderly offenders studied by Needham-Bennett *et al.*¹², 97 (65%) were cautioned and in 42 (28%) there was no further action. The police, however, referred only half of those later identified as psychiatric cases to welfare agencies, suggesting that closer links between the police and community psychiatric teams for the elderly might facilitate the identification of unmet mental health needs at an early stage.

The elderly serving sentences comprise both those imprisoned for the first time in old age and those who have grown old in prison. About 1% of the prison population in the UK is aged 60 + and about 5% in the USA. The number of sentenced prisoners in the UK aged 60 + has increased in recent years, from 333 in 1988 to 1055 in 1998, although there has not been a similar increase in the elderly remand population¹. A case history-based study in the USA of 25 new elderly offenders, most of whom were imprisoned for sex offences or homicide¹⁹, drew attention to their initial reaction to imprisonment, often being characterized by family conflict, depression, suicidal thoughts and a fear of dying in prison¹⁹. The physical health of older prisoners is also important, with about half of older prisoners having a long-standing illness or disability²⁰. The prison system makes scant provision for vulnerable groups and older prisoners may experience particular difficulties in adjustment on release.

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Sleep and Ageing: Disorders and Management

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Sleep complaints are very common in the elderly, with up to 40% being affected by insomnia. Further, a disproportionately large number of prescriptions of sedative-hypnotics are given to elderly people. Sleep disturbances in the elderly may be the result of physiological changes with the ageing process, poor sleep hygiene, medical and psychiatric conditions (particularly depression) leading to secondary sleep disturbance, and primary sleep disorders¹.

A number of changes in sleep characteristics and architecture occurs with ageing. In general, the older person takes more time to fall asleep, has more awakenings and less efficient sleep, as well as more napping in the daytime. Moreover, the older person tends to go to bed early and rise early, reflecting a phase-advanced rhythm of the sleep-wake pattern. In addition, there is a decrease in slow wave sleep and rapid eye movement (REM) sleep but an increase in light sleep, i.e. stages 1 and 2 of NREM sleep¹⁻³.

Inadequate sleep hygiene includes poor sleep habits and engaging in sleep-incompatible behaviour⁴. Excessive time in bed, irregular hour of going to bed, lack of exercise, excessive caffeine intake, alcohol withdrawal and noisy environment are all factors that might influence sleep.

Numerous medical conditions can lead to sleep disturbances, especially when pain is a significant feature. In the elderly, dementia is a common cause of sleep disturbance. In Alzheimer's disease, there is a decrease in slow-wave sleep and REM sleep, with increased fragmentation of sleep. A disrupted sleep-wake cycle is frequently found. Agitation and confusion in the evening and at night (sundowning) may also occur in some patients⁵.

As for primary sleep disorders, sleep apnoea, REM sleep behaviour disorder and periodic leg movement during sleep (PLMS) are the ones with increased prevalence in the elderly, and will be dealt with in this chapter.

SLEEP APNOEA

A period of apnoea is defined by a cessation of breathing for 10 s or more, whereas a hypopnoea period is a 50% reduction in the respiratory depth for 10 seconds or more. Sleep apnoea is characterized by recurrent episodes of apnoea and hypopnoea during sleep, and is usually associated with oxygen desaturation in the blood⁶.

There are two main types of sleep apnoea, obstructive and central. Obstructive sleep apnoea is the more common form. Cardinal features are loud snoring and excessive daytime sleepiness. Associated features include headache, insomnia, apnoea observed during sleep, excessive movements during

sleep, cognitive impairment, personality changes and enuresis. Physical problems include systemic hypertension, pulmonary hypertension and cor pulmonale⁶, which might explain why sleep apnoea is associated with an increased mortality due to cardiovascular events.

The exact prevalence of sleep apnoea is unknown but it is estimated that about 2-4% of the general population meet minimal criteria for obstructive sleep apnoea⁷. Its frequency increases with age, reaching a maximum between 50 and 70 years of age, and there is a male predominance. In the elderly, prevalence rates of 26-73% have been reported in various studies⁸. This shows that disordered breathing is a common problem in the elderly. However, a major unresolved issue is whether sleep apnoea is a less pathological condition in the elderly. Studies in clinical populations have shown that disturbed respiration during sleep in the elderly has minimal association with mortality and morbidity, while epidemiologic studies in the elderly have suggested otherwise². Pending further studies to clarify the issue, older people with symptomatic sleep apnoea probably should be treated in the same way as younger patients.

In the management of sleep apnoea, general measures include weight reduction and avoidance of alcohol and benzodiazepines before bedtime. In the majority of patients with sleep apnoea, continuous positive airway pressure (CPAP) during sleep is the treatment of choice. Surgical treatment may be indicated for patients with specific upper airway abnormality who have failed CPAP therapy or did not want CPAP for various reasons, such as frequent travelling⁹.

REM SLEEP BEHAVIOUR DISORDER (RSD)

This is a recently described parasomnia¹⁰. Presenting features are usually excessive motor activity during sleep, which may lead to repeated injuries to the patients or their bed-partners. Patients may talk or shout aloud in sleep, accompanied by vigorous limb movements, walking, falling out of bed, or carrying out various activities in their dreams. After awakening, patients may recall dreams that coincide with their motor activities. This suggests that the motor activity in sleep is a form of dream enactment. Some patients resort to various measures to protect themselves from injury, like tying themselves to the bed or putting a mattress on the floor. In addition, the nature of their dreams may change over the years, becoming very vivid and action-packed¹¹. The diagnosis of RSD should be considered in elderly people presenting with sleep-related injury or violence.

There are very few studies on the prevalence of RSBD. A study in the general population by telephone interview found that 0.5% had probable RSBD¹². Another study on a community sample of elderly reported a rate of 0.4%¹³. RSBD tends to occur predominantly in the elderly, and males are more affected. In general, the awareness of this condition is low among the public as well as clinicians, and misdiagnosis is common.

RSBD occurs in transient or chronic form¹¹. The transient form may be induced by drugs as well as alcohol withdrawal. The chronic form is either idiopathic or associated with neurological disorders in up to 50% of cases, such as dementia, Parkinson's disease, multiple system atrophy and vascular or neoplastic lesions of the central nervous system. In particular, there is a strong association with Parkinson's disease and dementia with Lewy bodies^{14,15}.

The exact aetiology and pathophysiology of RSBD is unclear. In normal people, there is generalized muscle paralysis during REM sleep, sparing only the diaphragm and extraocular muscles. In RSBD there is a disruption of this pattern and patients can thus move and act out their dreams¹¹.

Management of RSBD includes drug treatment with clonazepam, which is effective in up to 90% of cases, as well as safety measures to protect the patients from injuries¹⁶.

PERIODIC LEG MOVEMENT IN SLEEP

Previously known as nocturnal myoclonus, PLMS consists of stereotyped, periodic jerky movements of the lower limbs, usually occurring in light sleep. Symptoms include leg jerks, insomnia, daytime sleepiness and sometimes cold feet¹⁷. PLMS is associated with restless legs syndrome (RLS), which is characterized by the presence of unpleasant sensation in the lower limbs occurring when the patient lies down in bed, frequently leading to insomnia. However, PLMS can occur independently of RLS.

PLMS may occur as an isolated finding, but has been observed in a number of pathological conditions, such as chronic myelopathies and peripheral neuropathies, uraemia and sleep apnoea¹⁸.

PLMS is very common in the elderly; studies have reported rates up to 45%¹⁷. Nevertheless, the relationship of PLMS and insomnia is still a matter of debate. It has been suggested that PLMS may be coincidental with sleep-wake disorders, rather than being a cause of it; indeed, many elderly with PLMS are completely asymptomatic.

Mild cases of PLMS may not need any treatment. For more severe cases, options of drug treatment include benzodiazepines (particularly clonazepam), L-dopa, bromocriptine and opiates¹⁸.

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Rating Scale for Aggressive Behaviour in the Elderly

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Aggressive behaviour is one of the commonest causes for psychogeriatric admissions. It is highly correlated with carers' distress and rejection, as well as a decision for long-term institutional care¹⁻². To rationally evaluate the effectiveness of intervention strategies for aggressive behaviour, it would be very important to identify suitable assessment tools. The Rating Scale for Aggressive Behavior in the Elderly (RAGE) is specifically designed for the assessment of this aspect of behavioural problems in psychogeriatric institutional settings³.

The RAGE is a 21-item rating scale administered by trained professionals, usually the nursing staff in the setting. The items include different dimensions of aggression (verbal, agitation and physical aggression), with no pre-defined diagnostic specifications. Aggressive behaviour is defined in the RAGE as "an overt act,

involving the delivery of noxious stimuli to another organism, object or self, which is clearly not accidental". The scale is observer-rated and the staff is requested to complete the questionnaire based on observation in the ward over a 3 day period.

Patel and Hope¹ reported that the RAGE demonstrated good psychometric properties, with satisfactory reliability and validity. Comparison of RAGE with two other commonly used scales for aggression, the Cohen–Mansfield Agitation Inventory and the Brief Agitation Inventory, revealed that the three scales all highly intercorrelated, with meaningful constructs⁴.

Studies using RAGE to measure aggressive behaviour in institutional settings revealed that about half of the elderly in psychogeriatric wards had positive ratings in aggressive episodes over a 3 day observation period. It was found that elderly schizophrenic patients, when compared with the demented elderly, were more frequently rated as overall aggressive, albeit with the same total scores in the RAGE⁵. The scores on the RAGE were found to be higher in moderate degree of dementia, and were also reported to be associated with activity disturbance and the presence of psychotic features⁶.

Table 1. Items in the RAGE

Item	Ratings
Demanding	0–3
Shouted	0–3
Swore	0–3
Disobeyed ward rules	0–3
Uncooperative	0–3
Irritable	0–3
Sarcastic	0–3
Impatient	0–3
Threatened to harm	0–3
Antisocial acts	0–3
Pushed others	0–3
Destroyed property	0–3
Angry with self	0–3
Attempted to kick	0–3
Attempted to hit	0–3
Attempted to bite	0–3
Used object to hurt others	0–3
Self-inflicted injury	0–3
Injury to others	0–3
Sedation or restraint	0–3
Overall aggressiveness	0–3

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Sexual Disorders

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THE CAUSE OF ERECTILE FAILURE

Although levels of free testosterone fall with age, this does not relate to levels of sexual activity^{1,2}, which correlate closely with age itself. Three hypotheses for the decline remain (although whether loss of interest leads to erectile failure or vice versa is in doubt); these are: (a) that increasing atheroma causes a failure of vasodilation sufficient to cause an erection; (b) that degenerative changes in the autonomic system are the cause; and (c) that levels of self-confidence decline with age, leading to psychogenic failure. Clinical impressions do not support the latter hypothesis, and a study of 28 men with erectile failure over age 65 compared to 25 aged-matched men who were potent showed no differences in their GHQ³.

Feldman *et al.*⁴ compared measures of health in 1290 males aged 40–70 in Massachusetts and found erectile failure to be increased with heart disease, diabetes, hypertension, depression and anger, and decreased by high-density cholesterol and by levels of dihydroepiandrosterone, a breakdown product of testosterone, but not by levels of testosterone itself. They did not measure neuronal ageing. Rowland *et al.*⁵ found that penile sensitivity was related to erectile response. In our study³ reported above, we found that only a measure of autonomic neuronal integrity (pilocarpine-induced sweating) distinguished the two groups fully. This might suggest that most elderly men with erectile failure would respond to corporeal vasodilators, which would be less effective if the cause was atheroma. The subject is well covered by Schiavi⁶.

TREATMENT

Counselling allows the couple to express their fears and inhibitions, whilst the therapist can educate them about the normal changes of ageing. Elderly men, for example, often overestimate the importance of penetration, compared to petting, in the pleasure they give to their partner. Physical remedies are

more widely used and include vacuum pumps⁷, intracavernosal⁸ and intraurethral⁹ administration of alprostadil, oral sildenafil¹⁰, which prolongs the action of cavernosal nitric oxide, and oxpentifylline for vasculogenic failure. Dopaminergic drugs can restore libido, if fear of prostatic cancer contraindicates testosterone. Disorders of female arousal are reduced by HRT and also respond to sildenafil¹¹.

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Phenomenology of Wandering

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With current increases in the elderly population, dementia and its related behavioral disturbances are receiving much more attention. Wandering is one of the common, often treatment-resistant behavioral concomitants of progressive dementia. Most research on wandering has focused on nursing home- or dementia clinic-based samples. In these samples, wandering occurs in up to 65% of patients at some point in the disease process¹⁻⁷.

DEFINITION AND PHENOMENOLOGY

There has been much debate about the appropriate definition of wandering, as well as its phenomenology. A major difficulty in wandering research has been the lack of a uniform definition and of reliable assessment instruments. Wandering was initially defined as “aimless movement”^{8,9}. More recent definitions have viewed wandering not only as aimless, but also as having purposeful intent^{10,11}. Wandering has further been classified, on the basis of phenomenology, as benign or problematic⁸. A benign wanderer is a patient who roams aimlessly and is easily redirectable, whereas a problem wanderer is one who is disruptive to family, staff or other residents and includes the individual who is resistant to redirection. Attempts have made to quantify wandering behavior into two broad categories, continuous and sporadic. “Continuous wanderers” are defined as ambulating more than 50% of their wakeful time, while “sporadic wanderers” move about for less than half of their wakeful time.

A common theme that is consistently supported in the literature is that wandering is primarily influenced by a continuity of behaviors from earlier premorbid times¹². Recent research¹³ has also focused on the relationship of premorbid personality and wandering. Overall, wandering is viewed as having a beneficial effect for the wanderer by fulfilling a particular need. De Leon *et al.*¹⁸ showed that wanderers had poorer parietal lobe functioning than subjects with similar degrees of cognitive impairment. However, in a multisite, random sample of 163 ambulatory, cognitively impaired subjects, wanderers showed significantly greater impairment in basic skills (orientation, memory and concentration) and in the higher-order skills of language, abstract thinking, judgement and spatial skills¹⁴. In Monsour and Robb’s¹² sample of wanderers, 36/44 subjects had dementia or Alzheimer’s disease; eight had cerebral vascular accidents or arteriosclerosis. Although wanderers may have more cognitive decline, they often exhibit an intact social facade, which masks their deficits¹⁵. Snyder *et al.* also identifies that wanderers had a higher number of psychosocial needs than non-wanderers on the Human Development Inventory (HDI).

Table 115.1 Why patients wander

Trying to find “home”
Trying to find bathroom
Are hungry
Are in pain
Acute medical or environmental trigger, stress, or loss
Drug side effect or withdrawal
Are bored

Triggers of Wandering

Table 115.1 lists some common reasons why patients wander. They may be trying to find “home” or their room, whereas some are merely driven by the urge to void and are trying to find a bathroom. Some patients may be scavenging for food because they are hungry. Wandering behavior may be due to pain, e.g. arthritis or other painful medical states, which the patient cannot communicate. If the wandering behavior is of abrupt onset, one needs to ask if an acute stress or loss, whether a medical (e.g. an infection) or environmental (e.g. death of a roommate or family member) factor has triggered the wandering behavior. At times, wandering may be due to boredom.

TREATMENT

Table 115.2 highlights general treatment approaches to wandering behavior. The management of wandering behavior can be classified into three groups according to the potential etiology of the wandering behavior: (a) medical; (b) psychosocial; and (c) environmental.

The most important cause of wandering to exclude is an acute or chronic underlying medical problem, for example, an abrupt arrhythmia. Also, could the need to be on the move be a side effect of medications? Akathisia with neuroleptics or agitation with

Table 115.2 Treatment of wandering: general approaches

Eliminate physical/chemical restraints
Wander-safe indoor and outdoor areas
Eliminate distracting light and noise from environment
Use of electronic elopement prevention devices and patient identification bracelets
Proper staffing
Staff education and training
Philosophy of rehabilitation

fluoxetine are examples. Perhaps a withdrawal syndrome, such as from alcohol or benzodiazepines, may be implicated. Pain as a trigger for wandering always needs to be considered.

In the nursing home, proper staffing, appropriate staff training with regard to causes and management of wandering behavior, coupled with supervised structured activities, such as music therapy and simple crafts, are important. Use of planned activities and distraction techniques may be helpful.

One of the key elements in managing wandering behaviors is to provide an appropriate and safe environment where wandering behavior can be tolerated. This should ideally include specially designed indoor and outdoor roaming areas that lack safety hazards or distracting elements. Innovative ideas include a "reduced stimulation unit"^{11,16} and a "wanderer's lounge program"¹⁷. It is vital that wandering behavior is not dealt with via the use of physical restraints or the heavy use of sedatives or tranquilizers. Patients often fight such measures, resulting in more significant problem behaviors.

A major concern for family or institutional caregivers is that the wandering patient may elope and come to physical harm, e.g. run into automobile traffic. The use of "Wander-Guard"-type devices on doors and a special identification bracelet, labeled "cognitively impaired—if lost, please call (phone number of facility)" may help.

CONCLUSIONS

Wandering exacts a heavy toll on family and professional caregivers. For the wanderer, issues of safety are of paramount concern. To ease the burden on caregivers, it is important to try to ascertain *why* the patient is wandering. This will usually give clues to appropriate management. Unfortunately, wandering patients usually cannot communicate why they are wandering because of their cognitive impairment. Consequently, caregivers need to place themselves in the patient's shoes to come up with the answers.

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Part L

The Presentation of Mental Illness in Elderly Persons in Different Cultures

Problems of Assessing Psychiatric Symptoms and Illness in Different Cultures

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Increasingly, psychiatrists must have the skills to assess and deliver care for people from different cultures. Many areas can only be touched on here, and several of the references stem from the cross-cultural literature in younger adults. The context is very broad, covering industrialized countries with multicultural populations and less-industrialized countries. It is worth noting that, due to a range of barriers, many people from different cultures have, of course, very poor access to psychiatric assessment and care¹⁻⁴.

COMMUNICATION AND CULTURAL AWARENESS

While health staff often come from different cultures than their patients⁵, they are usually working within at least a familiar framework and landscape. With someone from a very different culture, the ability to assess appearance, behaviour and symptoms will be limited. The practitioner must work harder to gather the information to make an accurate assessment and to develop rapport and trust. Awareness of one's lack of information of one's own cultural "encumbrances" and of potential strong feelings about each other's cultures is crucial⁶. Training and consultation with a wide circle of people from the relevant culture is needed. Ask the patient early on about his/her background and, for migrants, his/her place of origin and experience in the new setting. Listening, asking open questions, acknowledging family expectations and a willingness to discuss issues such as racism and social needs are all vital. Within reasonable professional boundaries, the psychiatrist should be willing to respond to some questions about him/herself. For example, with older Jamaican migrants, culture-specific care providers stressed that making "a connection" with the doctor would facilitate the assessment and acceptance of care⁷. Show sensitivity but do not neglect important areas on "cultural" grounds, such as alcohol intake in Muslims. Apparent persecutory ideas must be explored and may reflect an appropriate response to injustice.

Language barriers should be addressed through the employment of ethnically close bilingual workers, otherwise use a competent interpreting and advocacy service⁸. Using relatives or other staff to interpret should be limited to emergencies. Requiring patients to speak in their second language can distort the clinical picture⁶. Aspects of non-verbal communication may

also differ, such as avoidance of direct eye-to-eye contact in some Asian and Pacific cultures.

DIFFERENT PRESENTATIONS OF EMOTIONAL DISTRESS

People from all cultures experience both somatic and psychological symptoms when emotionally distressed⁹. One reason for a more somatic presentation may be that this is seen as a more appropriate focus for medical consultation by some cultures. Second, there may be a continuum of experience and interpretation, with the more "somatizing" cultures at one end and the more "psychologizing"^{10,11} at the other. Thirdly, a large number of "somatic" complaints are actually metaphors for mental distress. Many of these relate to heart discomfort (e.g. a heart that is "sinking" or "uncomfortable")¹²⁻¹⁴ and to abdominal sensation^{15,16}.

Culture-bound syndromes usually represent cultural explanations for recognizable psychoses or neuroses¹³. For example, Dhat, a belief that semen is leaking from the body in urine, is a complaint in India. It may be a presenting feature, and is used as an explanation for weakness due to depression or organic disease.

DEPRESSION

There has been much debate about the existence of depression as a universal cross-cultural category^{10,17}. It is reasonably established that depressive disorders exist across cultures and are strongly related to local constructs^{15,18,19}. However, symptoms vary (e.g. depressed older Jamaicans and African-Americans describe feeling "low", "bad" or "fed-up")^{20,21}; multiple somatic symptoms or metaphors may be presented; and some cultures may emphasize their explanation (e.g. social or spiritual) for their symptoms²². The validity of screening scales may vary, e.g. a lower cut-off point for the Geriatric Depression Scale has been recommended for older African-Caribbeans, African-Americans and Mexican-Americans^{23,24}. Symptom profiles in depressed cases may also differ considerably across cultures, as shown even across European centres in older people³⁴. Depression at community level also appears to be highly correlated with anxiety²⁵. The implications are that: (a) it is important to enquire about the *full* range of affective and neurotic symptoms; and (b) a wide definition of mood disorders is likely to be most useful in the clinical setting.

FUNCTIONAL PSYCHOSES

Psychotic symptoms and signs appear similar in form across a wide variety of settings²⁶, although the content, e.g. of delusions, will be influenced by culture. Also, of course, the patient and his family's explanation for the illness will depend on their cultural framework²⁷. Hence, someone from Zambia may explain his/her schizophrenia as he/she might explain a stroke, depression or a burglary—as due to bewitchment or to having angered a spirit.

Culturally-supported dissociative states and altered states of consciousness can be misleading⁶. Pseudohallucinatory phenomena have been described in non-psychotic depression, anxiety and distress states^{13,28}.

An apparently unusual idea, such as believing oneself to be bewitched, is only a delusion if it is out of keeping with the beliefs of others in the culture. This must be checked by asking someone with appropriate knowledge. If faced with the patient alone, ask how he/she came to believe this, and if others close to him/her agree. If a traditional healer told him/her so, and his/her peers agree, then it is at least likely that this is a culturally sanctioned belief. Another error is to assume that a belief is culturally normal when it is actually abnormal.

DEMENTIA

DSM-IV criteria²⁹ for dementia require the demonstration of cognitive impairment of sufficient severity to interfere with the activities of daily living. However, cultures vary in the extent to which they expect older people to take responsibility, e.g. for domestic activities. Also, impairment from physical conditions is commoner in socially disadvantaged people and will be difficult to distinguish from that due to dementia, requiring greater emphasis on physical examination and tests^{6,30}. Many cognitive tests include items affected by education (e.g. requiring reading, writing or arithmetic skills) and/or which may have little relevance in certain cultures (e.g. "Who is the President?", "Take 7 from 100"). When testing those unfamiliar with such approaches, be courteous, encouraging (without "helping"), give explicit instructions and some dummy tasks to allay anxiety³⁰. Rather than doggedly adhering to the original version of instruments, it is appropriate either to make rational adaptations (as described for the MMSE³⁵) or to develop new instruments³¹. Novel adaptations will, of course, require translation, back-translation and pre-testing³³. Suitable informants should be screened for any decline in function of their close contact³². The Community Screening Instrument for Dementia^{31,33} combines culture-fair cognitive testing with a structured informant interview. This approach should become more routinely applied, both in primary and secondary care.

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Depression in the Indian Subcontinent

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Common mental disorders, such as depression in late life, are not well understood or acknowledged by either the community or the medical profession in India, for three important reasons. First, recognition of common mental disorders as a psychiatric problem in general is relatively poor. Most patients present with somatic symptoms. While many patients and health providers acknowledge the non-organic basis of these symptoms, neither group is comfortable with labels that imply a relationship to psychiatry. Indeed, terms such as “depression” and “anxiety”, when used to define diagnostic categories of psychiatric disorder, have no conceptual equivalent in any Indian language. Ethnographic studies have shown that symptoms of depression are attributed to tension, family conflict and lack of family affection, rather than being seen as a biomedical psychiatric problem¹. The second reason for low recognition is that the relative proportion of elders is less than 5% for most Indian communities. This is bound to change in the future, with falling birth-rates and rising longevity leading to predictions that, over 20 years, this oldest sector of the population will exceed 100 million. The implications are grave, for India has no systematic social welfare system for the aged, and is faced with the gradual breakdown of traditional extended family systems that have formed the bulwark for the care of the disabled and chronically ill¹. The third reason is:

the fatalistic attitude toward aging in India, which mandates that elderly persons accept their physical and mental condition as a normal part of old age. Not only are elderly persons with mental illnesses rarely brought to a physician, but those with treatable medical conditions also often receive no medical attention².

There are few epidemiological studies of common mental disorders in elders in India and no published studies, to date, that have used structured psychiatric interviews. Prevalence rates for depression in a community sample of elders have varied from 6% in southern India³ to over 50% in rural West Bengal⁴. The common presenting complaints are tiredness, sleep complaints, aches, tingling-numbness in the hands and palpitations. On enquiry, however, most depressed elders will admit to cognitive and emotional symptoms typical of depression. The hallmark cognitive feature is anhedonia, or loss of interest. Suicidal feelings and agitation are also common³. The suicide rate in the 50+ age group (12/100 000) is nearly twice the national average (7/100 000). Co-morbidity with physical ill-health is common; by some estimates, more than 90% of elders with a psychiatric disorder also have some physical disorder³. Risk factors for depression include low education, poverty, social isolation and family discord.

The latter is on the rise as a result of the breakdown of traditional community structures resulting from the massive migration of the younger productive members of families to urban areas and reduced economic activity in rural areas. The commonest treatments in primary care are symptomatic. Thus, benzodiazepines for insomnia and vitamins and “tonics” for tiredness are amongst the commonest prescriptions for common mental disorders in general health care, while antidepressants or psychotherapy are rarely offered⁵.

The rising rates of recognized risk factors, relatively low recognition of depression and even lower rates of appropriate interventions should cause considerable concern to public health policy and planning in India. One major limitation in influencing policy and practice is the lack of systematic evidence of the epidemiology of depression, and the efficacy and cost-effectiveness of treatments for depression, in elders in India. Research into the mental health needs of elders in India is clearly an important area for future psychiatric research. Health education should aim to educate health workers and the community to recognize that anhedonia and insomnia are not the expected price of growing old, but the result of a common, disabling and treatable illness. Removing stigma may require integrating the subject of depression into training programs for community and general health workers, and collaborating with non-governmental organizations that are pioneering programs to empower the elderly, support families with a mentally ill elder and provide health care sensitive to their needs. Working with the existing manpower and health and social service infrastructure is likely to be more successful in meeting the mental health needs of elders in India than developing specialized psychogeriatric services throughout the country.

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Dementia in the Indian Subcontinent

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Of India's population of nearly one billion, 50 million are aged 65 years or older and constitute a major potential high-risk group for dementia, yet pathological studies of dementia in India suggest that Alzheimer's disease (AD) is rare or unrecognized in most clinic populations. The few published epidemiological studies of dementia from India suggest potential regional differences in prevalence, which may be partly attributable to literacy or urban/rural residence. Differences across studies may also reflect methodological difficulties and differences. Psychometric and other screening instruments must be standardized in different Indian languages for elderly populations with widely varying levels of literacy, education and urbanization. Older individuals frequently have inadequately corrected sensory impairments, which can interfere with testing. They may have little interest in current national or world events, and thus appear to be impaired, and may not know their dates of birth. Cognitive deficits may be under-recognized and under-reported by family members, out of respect for, as well as reflecting low expectations of, the elderly. Prevalence rates of dementia from different studies in India¹⁻⁵ have ranged from a low of 1.36% to a high of 3.5% among those aged 65+. These low rates were found despite the use of highly standardized instruments for case detection, e.g. the GMS-AGECAT program and the community version of the GMS at the Madras site of a WHO multicenter study⁴.

Lower prevalence may be due to shorter life expectancy, with selective survival of those not at risk for dementia, and also to shorter duration or survival with the disease. Survival may be underestimated if the manifestations of dementia are detected late

or attributed in their earlier stages to normal aging. Plausible risk factors yet to be explored in Indian populations include head trauma, thyroid disease and illiteracy. Potential protective factors might range from family caregiving to low-fat diets. Tolerance of memory loss in old age, as well as lack of financial resources, may delay acceptance of treatment across the majority of Indian communities. Although prevalence is low, dementia may still pose a major public health challenge given the vastly growing population, the minimal existing infrastructure and the transitions in family structure in these regions.

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Dementia and Depression in Africa

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Africa is a multicultural, multiracial society, consisting mainly of Negroid people, although Arabs predominate in the north and white settlers constitute a minority in the south. There are no policies and programmes for old age in most countries of Africa, and the majority of elderly people live in often neglected rural areas. Most older Africans are impoverished, have little or no education and depend on their children for sustenance. Provision for their medical care is grossly inadequate. In this chapter the situation of the elderly in Nigeria will be used as a prototype for Africa.

Only about 3% of Nigerians are aged 65 years and older¹. The thrust of healthcare policy is towards communicable diseases in children, and geriatrics and geriatric psychiatry are both in their infancy. About the only policy document regarding old age care states that nursing homes will be discouraged, while home visits to older citizens will be encouraged and day care centres will be established. Federal, state and local governments are expected to share responsibilities for the care of older citizens²; however, as with most policies, there is a big gap between conception and implementation.

RESEARCH

Although geriatric research is limited, a number of studies have looked into dementia and depression. Because of the low level of education, most questionnaires designed in Western societies will require modification before application to older persons in Africa, especially in those that measure cognitive functions. Role expectations of older persons are also different, e.g. they are not expected to do household chores in the multigenerational living arrangements which are popular.

Dementia

Initial publications on dementia in Africa were about hospital patients, Lambo³ in Nigeria and Ben-Arie *et al.*⁴ in coloured older persons in South Africa. An earlier community study on dementia created the impression that Alzheimer's disease (AD) was rare or non-existent in Nigerians⁵. This conclusion is probably related to methodological issues in the study.

A major community-based study comparing older African-Americans in Indianapolis with older Nigerians in Ibadan has been ongoing since 1992. Prevalence rates of both dementia and AD were significantly higher in Indianapolis compared with Ibadan, 4.82% vs. 2.29% for dementia and 3.69% vs. 1.41% for AD⁶. More importantly, there was a progressive increase in the

prevalence rates of both dementia and AD with increasing age after 65 years. Another important finding was that the Apoe4 allele, which has been reported to be associated with AD in most studies, was found to be unrelated to AD in Nigerians⁷. Important risk factors for dementia in Nigerians included age and female sex only; other well-known risk factors were not identified¹⁵. Behavioural disorder symptoms were found both in Nigerians and African-Americans but it was felt that Nigerians are generally more tolerant of behavioural disorder symptoms in their demented family members, who are often not treated⁸.

In another major community survey of people aged 60+ in Egypt, Farrag *et al.*⁹ reported a prevalence rate of 4.5% for all dementias, 2.2% for AD and 0.9% for multi-infarct dementia. In that study also the rate of dementia doubled every 5 years.

Depression

Lambo³ diagnosed more depression in hospital older male patients compared to older females; Baiyewu *et al.*¹⁰ reported that 5.4% of older Nigerians were depressed in a community survey with a preponderance of males. Recently, Sokoya¹¹ also showed that 5.4% of older primary care attendees were depressed, using AGE-CAT criteria¹². Depression was related to low income but there was no gender difference. Depression was not often recognized and treated.

Nursing Homes

Nursing homes are very few in Nigeria, as in other African countries. However, in a study of psychiatric disorders in two Lagos nursing homes, using AGE-CAT and DSM-III-R¹³ diagnoses, 48% of patients had dementia and 17% had depression¹⁴. Although the sample size was small and it is difficult to generalize, the figures are close to those reported in centres in Western societies, which may portend that the problem will become more evident in future as the number of older persons increases.

FUTURE TRENDS

There have been two major community studies on dementia in Africa; the study among the Arabs of Egypt gave prevalence rates twice as high as for Nigerians. There is an urgent need to have more studies on psychiatric disorders among older persons in Africa. Such studies will be informative on rates, patterns of illness and risk factors, as well as assisting in policy formulation.

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Mental Illness in South America

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This chapter will present some features of psychogeriatrics in Latin America. With this aim in mind, the situation with regard to epidemiology, risk factors, recognition of symptoms, the influence of culture and the state of development of services will be outlined.

EPIDEMIOLOGY

Depression

Estimates of the frequency of depression symptoms vary widely from 8 to 46.2% (see Table 118.1). Such a variation depends heavily on the instruments used, cut-off points and severity of the symptoms. Using higher cut-off points, Cornejo and Lazlo¹ obtained a prevalence of 8%. Eisirik², using the DSM-III-R checklist, found a prevalence of 4.7% for major depression in an urban sample. Women presented higher rates, 5.8% vs. 2% for men.

Dementia

The prevalence of organic brain syndromes varies (4.3–29.7%). The study of Veras and Coutinho³ investigated three subdistricts of the city of Rio de Janeiro. The figures varied dramatically: 5.9 in Copacabana subdistrict to 29.7 in Santa Cruz. Several socioeconomic variables accounted for such differences.

Cognitive impairment ranged from 10.5% to 29%. The study conducted by Xavier⁴ relied on the oldest sample (80+) and investigated the age-associated cognitive decline through a neuropsychological battery (19.7%). Silberman *et al.*⁵ have found the highest figure (29%) in a small urban community sample.

Herrera *et al.*⁶ conducted a survey using a three-stage design in the city of Catanduva in the state of São Paulo. The MMSE was used as a screening instrument. All the suspected cases were further investigated by a specialist, using a set of evaluations. All subjects with a diagnosis of dementia went through a complete laboratory examination; 1660 subjects aged 65+ were interviewed. The prevalence of dementia was 7.1%. In the series of 118 cases, as many as 64 cases (54.1%) were diagnosed as having Alzheimer's disease, 11 (9.3%) vascular dementia and 17 (14.4%) an association of vascular dementia and Alzheimer's disease. The prevalence of dementia was age-associated: 1.3% in the age group 65–69; 3.4%, 70–74; 6.7%, 75–79; 17%, 80–84; and 36.9%, 85+. Education was an important protective variable against dementia. The prevalence rates can be seen in Table 118.2.

Table 118.1 Prevalence of depression symptoms in Latin America

Ref.	Age	Instrument	Sample size	Prevalence (%)
16	65+	Zung SRDS	26	46.2
1	68+	Brink GDS	433	8 (severe depression)
3	60+	Short care	738	20.9–36.8
5	60+	MADRAS	62	30
2	60+	MADRAS	344	21.5
—		DSM-III-R		4.7 (major depression)

Prevalence of Mental Disorders

Ramos *et al.*⁷ conducted a household survey using a multidimensional functional assessment questionnaire (OARS methodology); 1062 elderly aged 60+ living in three subdistricts of the city of São Paulo, Brazil, were interviewed. Among other data 27% of the subjects were considered psychiatric cases, as assessed by the mental health screening questionnaire included in the methodology. Eisirik², using the SRQ as a screening instrument, observed that 10.2% of subjects were considered positive psychiatric cases in a sample of 344 individuals aged 60+.

Alcoholism

Community studies are mainly focused on adult samples. Data concerning elderly populations are usually restricted to a small number of elderly persons. Yamamoto *et al.*⁸, studying an unselected sample in Lima, Peru ($n=815$), screened seven men (43.75%) out of 29 subjects aged 65+ for lifetime alcohol abuse or dependence.

RISK FACTORS

The Latin-American literature is not extensive in this respect. Epidemiological data⁶ have indicated that dementia of the Alzheimer's type occurs somewhat more often in women, in a proportion of 2:1. Sadigursky and Oliveira⁹ investigated the association between religious practice and depression on women aged 60+ attending a geriatric clinic. This case-controlled study examined 90 women. They found that the regular practice of religious activities had a protective effect on depression (7.4%) compared with less interested in religious activities (44.4%).

Table 118.2 Prevalence of dementia in Latin America

Reference	Age	Instrument	Sample size	Prevalence (%)
Organic brain syndromes				
18	55+	QMPA	139	4.3
	65+		44	6.8
12	65+	FHT	91	5.5
3	60+	Short care	738	5.9–29.7
Cognitive impairment				
5	60+	MMSE	62	29
2	60+	MMSE	344	24.7
4	80+	Battery	77	19.7
17	60+			10.5
Dementia				
6	65+	MMSE	1660	7.1

Studies of adaptation to high altitudes have led investigators to examine whether altitude itself is associated with mental problems in late life. Countries like Peru or Bolivia have a peculiar geography suitable for these type of studies. They have communities living both at sea level and at 4000 m above sea level. Alarcón *et al.*¹⁰ examined 93 subjects aged 60+ living in Lima (150 m) and 140 in Cusco (3400 m). Among other findings, they verified that the frequency of symptoms of anxiety and cognitive decline was greater at high altitudes than at sea-level.

RECOGNITION OF SYMPTOMS AND THE INFLUENCE OF CULTURE

An ethnographic study conducted in north-western Brazil examined the meaning of sadness in this low-income population. This qualitative study, which included retired subjects aged 60 years, found three main semantic clusters for sadness: an inner set, the body set and the interaction set¹¹.

Cross-cultural comparisons between unselected samples of elderly in two urban populations, Mannheim, Germany, and São Paulo, Brazil, using similar instruments, found similar symptom profiles in both cities. Some disparities between the samples are thought to be real and to relate to sociocultural differences as well as the greater stresses of daily life in São Paulo¹².

STATE OF THE DEVELOPMENT OF SERVICES

The pattern of service utilization in Latin-America by older patients with mental health problems is not well known. Few studies have investigated the ability of primary care services to meet the demand¹³ and to determine different potential users' expectations and needs¹⁴. It is reasonable to suppose that important inter-regional differences exist in the delivery of care. In some

places the only source of available psychiatric treatment is the hospital, generating a misuse of psychiatric resources¹⁵.

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Part M

The Practice of Psychogeriatric Medicine

- MI The British Model of the Organization of Services
- MII The North American Model
of the Organization of Services
- MIII Liaison with Medical and Surgical Teams
- MIV Rehabilitation and General Care
- MV Prevention of the Mental Disorders of Old Age
- MVI Education

Psychiatry of the Elderly—the WPA/WHO Consensus Statements

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Between 1997 and 1999, under the Chairmanship of the late Professor Jean Wertheimer, the World Psychiatric Association, Section of Geriatric Psychiatry, published a series of consensus statements in collaboration with the World Health Organization Division of Mental Health. A wide range of non-governmental organizations (NGOs) participated.

The first Consensus Statement¹ defined the scope of the specialty, emphasizing that this has become necessary because of increasing longevity (most markedly in the developing world); the relatively high prevalence of both functional mental disorders and the dementias in old age; and the need to adopt a multidisciplinary approach, while defining professional roles within the team. The scope of psychiatry of the elderly is “the psychiatry of ‘retired’ people”, and includes the full range of mental illnesses, including affective disorders, psychoses, substance abuse, the dementias and the mental health problems of “graduates” whose mental health problems continue into old age. The specialty also needs to address psychological, physical and social aspects of mental health problems in older people and the biosociocultural changes associated with ageing.

The characteristics of a successful psychiatry service for the elderly are summarized as community orientation, a multidisciplinary approach, an emphasis on abilities as well as deficits, and a core aim to improve quality of life rather than simply to alleviate symptoms. The main objectives of treatment are the restoration of health, improvement of quality of life, the minimization of disability, the preservation of autonomy, and addressing the needs of family and other carers as well as those of the individual patient. The high relapse rate of functional psychiatric problems in old age necessitates close follow-up after successful initial treatment.

Priorities within any new specialist service include teaching psychiatry of the elderly to primary healthcare workers; training existing mental health professionals in special mental health problems of the elderly, and establishing at least one multidisciplinary resource/expertise centre.

The second Consensus Statement² identified general principles that should underpin any quality specialist psychiatry of the elderly service:

- Good health and optimal quality of life are fundamental human rights irrespective of age or mental disorder
- All people have right of access to appropriate services
- Recognized needs should, within resource constraints, be met appropriately and ethically

- This can only be achieved through health and social measures adapted to local needs
- Older people with mental health problems and carers should be involved, individually and collectively, in care planning
- Governments should recognize the crucial role of non-governmental organisations and work with them.

The specific qualities of a good service are that it should be Comprehensive, Accessible, Responsive, Individualized, Transdisciplinary, Accountable and Systemic (C̄ARITAS), and should attempt to both prevent mental health problems from arising and identify them early when they do arise. As well as offering comprehensive assessment and acute management, the service should provide continuing care and support to patients and carers and address spiritual and leisure needs as well as medical needs.

The third Consensus Statement³ focused on education. Targets for educational initiatives include: health and social care professionals at undergraduate, postgraduate and continuing education levels; health and social service managers; other care workers; family and other informal carers; voluntary workers; public policy makers; and the general public. Although such a wide range of target groups inevitably start with different needs and different starting levels of knowledge, a generic core curriculum can be derived from the learning needs of health professionals. Essential curriculum elements include:

- Processes of ageing in individuals.
- The demography, economics and politics of ageing societies.
- The epidemiology, pathology, clinical features, assessment, diagnosis, treatment and management of the mental disorders of old age.
- Physical disorders and impairments of function that commonly occur in old age.
- The special significance in old age of the interdependence of mental, physical and social factors.
- Principles of health promotion and the preventive psychiatry of old age.
- Ethical and legal issues relevant to older people.
- Principles of planning, provision and evaluation of services in different settings.
- Needs of carers and approaches to their support.
- End-of-life issues.
- Principles and practice of multidisciplinary teamwork.
- Interviewing and communication skills.
- Fostering of positive attitudes and insight into the reasons for negative attitudes.

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Development of Health and Social Services in the UK in the Twentieth Century

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IMPERIAL BEGINNINGS: THE POOR LAW AND THE ASYLUM

In Britain, the twentieth century dawned in a blaze of imperial glory. Three years earlier, Queen Victoria's diamond jubilee had been celebrated across the globe with a splendid procession in London itself. The mood of the nation was confident, even optimistic, and world-domination was accepted almost as a birthright of the British people. Britain had survived the rigours of the Industrial Revolution and had come up fighting. Yet, at this time, public health measures were rudimentary and confined largely to the establishment (in 1848) of sanitary authorities with medical officers of health to oversee sewers and water supplies. The poor law was still in force and poor law institutions were made deliberately unpleasant. This followed the principle of "lesser eligibility", set out in 1834, which stated that those receiving poor law assistance should not be as "eligible" (well provided for) as an "independent labourer of the lowest class"¹. For the poor sick this had been ameliorated, to some extent, by the setting up of poor law infirmaries in 1868, but there was still a vast gulf between these institutions and the voluntary hospitals, which were supported by rich philanthropists. Retirement pensions, even retirement itself, were things of the future and there was an association between poverty, ill-health and old age which was recognized by an 1895 Royal Commission on the aged poor.

Mentally ill people were still incarcerated in large county asylums. In 1808, partly as a response to the appalling conditions in some private "madhouses", local magistrates had been given the power to set up asylums and in 1845 this provision had been made mandatory.

From 1900 onwards, developments have been influenced by major world events, political philosophy, public opinion and the power of pressure groups. The Boer War, starting in 1899, revealed the poor physical fitness and ill-health of many young men. Improvements in midwifery and child care were soon legislated for, with school meals starting in 1906 and the notification of live births, health visiting and the school medical service soon following. In 1908 the first national scheme for old age pensions was set up to try to alleviate poverty amongst old people. It was non-contributory and means-tested. Initially, recipients also had to be "of good character"

The Royal Commission on the Poor Laws and the Relief of Distress in 1909 considered most of the issues of domiciliary and hospital medical care. A minority report condemned the poor law institutions as a public scandal, with the infirmaries understaffed

and lacking skilled medical input². Out of hospital, the poor law doctors had no contact with local authority public health services, the voluntary dispensaries were overcrowded and ineffective and the medical clubs, financed by workers' subscriptions, underpaid their doctors and did not cater for the chronic sick or dependants. The writers of this report dismissed the idea of a medical insurance system.

Yet, in 1911, the establishment of such a system marked an important development in the evolution of general practice in the UK. The medical profession fought for, and won, independence and capitation fees rather than a salaried service, and administration by insurance-based panels rather than local authorities³. Higher income groups, families and hospital care were excluded but the scheme was nevertheless a qualified success.

THE MINISTRY OF HEALTH: BETWEEN THE WARS

In 1918 the Ministry of Health for England and Wales was formed and the Minister quickly appointed a consultative council, which in 1920 produced a report described by Pater² as "nothing less than the outline of a national health service" (p. 7). Their scheme might well have avoided some of the split between general practitioners and hospital doctors that has been one of the problems of the National Health Service (NHS) as it was eventually implemented.

Control of the workhouses passed to local authorities in 1930, the beginning of the end for the poor law. After a post-war cash crisis, the voluntary hospitals continued, becoming more specialized in acute care and leaving the chronic sick and infectious diseases to the local authorities. A number of reports pressed for a more coordinated hospital system and for universal health insurance. Knowledge was advancing. In 1935 Warren^{4,5} began her work in developing geriatric medicine and, a few years later, pioneers began to write of the issues concerning old people with mental illness⁶⁻⁸.

Before the Second World War, the Emergency Medical Service (EMS) was set up to cope with expected severe civilian casualties from the bombing of cities. On the declaration of war, 140 000 people, many of them elderly, were discharged from hospital over 2 days⁹. The EMS also coordinated the work of the voluntary and local authority hospitals, providing the framework for the future NHS Regional Hospital Boards. Physicians and surgeons from the elitist voluntary hospitals came face to face with the conditions of the poor law institutions.

THE POST-WAR NATIONAL HEALTH SERVICE

The last of the series of British Medical Association (BMA) reports pressing for reform in 1942 coincided with the Beveridge report and was followed in 1944 by the NHS White Paper, enacted in 1946 and effective in 1948.

The National Health Service, as then set up, was tripartite. Primary care services—general practitioners, opticians, dentists and pharmacists—were answerable to local executive committees; maternity, child welfare, health visiting, health education, immunization and ambulances remained the responsibility of the local authority; and hospitals were administered by Regional Hospital Boards with teaching hospitals retaining boards of governors directly answerable to the Ministry of Health. One of the assumptions when the NHS was set up was that increasing health in the population would cause health expenditure to level off. It never did, and in 1956 the Guillebaud Committee, appointed to find ways of avoiding a rising charge upon the exchequer, concluded that there was no evidence of inefficiency or extravagance in the NHS. In fact, the committee was concerned about a lack of capital expenditure (a concern again of relevance more recently). In 1962, this problem was addressed in the Hospital Plan.

Meanwhile, in the mental health field, the idea of community care was gaining ground. Tinker¹⁰ attributed this to five factors. First, there was a general dissatisfaction with institutional care and a search for alternatives. Some of the experiments in the “therapeutic community” work of the Second World War had challenged the accepted authoritarian culture of the mental hospital¹¹. In addition, the advent of electroconvulsive therapy (ECT), antipsychotics and effective antidepressants facilitated the move away from custodial care to medical treatment at home or in ordinary hospitals. Next, there were beginning to be practical problems in running residential establishments, including staff recruitment. Then there was concern about the cost of institutional care and, finally, a recognition that mentally ill people were entitled to live in as normal a way as permitted by modern treatments. The 1959 Mental Health Act liberalized the treatment of mentally ill people and opened the way for a move away from the old psychiatric hospitals to the new concept of psychiatric units attached to the district general hospitals of the 1962 Hospital Plan.

The large institutions were, in any case, rocked by a series of scandals about the mistreatment of patients. This resulted in the establishment of the Hospital Advisory Service (later the Health Advisory Service), effectively an inspectorate to monitor standards and spread good practice.

In general practice, a financial allowance for practices in deprived areas combined with other factors to promote the rapid development of local health centres and group practices from the mid-1960s. Local authorities produced their own health and welfare plans but there was poor coordination with the hospital authorities and the general practitioners’ executive committees. Within the local authorities, the Seebohm report (1968) was followed by the Social Services Act, which required the setting up of social services departments. The Department of Health and Social Security was created in 1968 by the amalgamation of the Ministries of Health and Social Security, a merger that lasted for some 20 years.

Reforms

In 1974, for the first time since its inception, the NHS itself was reorganized. The chief elements of this reorganization were the separation out of health and social services functions, the integration of all health functions under one management and

the establishment of area health authorities, generally co-terminous with local authorities, to facilitate joint planning. Community Health Councils were also created to represent the views of consumers. Unfortunately, the reformed service did not work well. There were too many layers of responsibility and taking decisions seemed to be delayed whilst information and responsibility were passed up and down the tree. There was an increase in clerical and administrative staff without a corresponding increase in managerial efficiency. During this period, important government reports were produced, including *Better Services for the Mentally Ill*¹² and *A Happier Old Age*¹³.

In 1982 the Area Health Authorities were abolished and new district health authorities combined the functions of the old areas and districts. In some areas, co-terminosity with local government was lost. A new government was determined to cut public expenditure and the rate of growth of the NHS slowed. Following the Griffiths report¹⁴, a general management structure was established within the NHS and Family Practitioner Committees became independent. Government payment for continuing care was channelled to the private sector and social services and hospital provision for this group of patients/residents was either reduced or failed to keep pace with demographic changes¹⁵.

Psychiatric services were coping with the implementation of the 1983 Mental Health Act, which set up time-consuming quasi-judicial procedures for reviewing patients who were detained in hospital under compulsory orders. The new Act also set up a Mental Health Act Commission to review treatment of detained patients and to advise on certain types of treatment.

MARKET FORCES: A RADICAL DEPARTURE?

Then came the most radical reform of the NHS attempted to that date, a reform not just of the service but of the basic philosophy of “service” underlying it. Some suspected that it was the beginning of the end for the National Health Service. The 1990 National Health Service and Community Care Act introduced the concept of an “Internal Market”. The new health authorities became planners and purchasers of health care at “arm’s length” from the providers, which were initially directly managed units (DMUs), and became semi-independent Trusts. The health authorities were provided with a budget for the local population and placed contracts for care with Trusts or the private and voluntary sector in order to obtain the best “value for money”. Quality was, at least in theory, specified in the contract and monitored.

Competition and other features of business life were “introduced” into the NHS, not least by setting up groups of “fundholding” general practitioners, who were enabled to make their own contracts for secondary care. Some of the changes were potentially positive, such as the setting up of Trust Boards to manage local services and an emphasis on sound financial regulation through corporate governance. Unfortunately, the bottom line was very clearly financial and in many cases clinical services were sacrificed to balance the books.

These proposals were pushed through in the teeth of strong opposition from staff and groups representing the consumer. Honigsbaum¹⁶ analysed the situation in 1990 and concluded that if patient care suffered, then “the nation may decide that the restraints imposed are not worth the savings they produce. Today, as in 1911 and 1948, it is the public interest that will predominate”. The medical profession were excluded from the plans for this reorganization. Klein concluded that, if a new political settlement were not reached between the government and the profession, it seemed unlikely that the NHS would survive long into the twenty-first century¹⁷.

A NEW NHS?

In fact, perhaps partly because of public dissatisfaction with what was happening to the NHS, the government was not re-elected and a radical reforming Labour government came to power. The engines of privatization and the internal market were reversed and new reforms were produced. In December 1997 a White Paper, *The new NHS: Modern, Dependable*¹⁸, outlined a comprehensive new vision for the NHS. Two of the main planks of the new policy were the setting up of primary care groups (PCGs) to replace the fundholding/non-fundholding split and the introduction of comprehensive quality controls to ensure high standards and equity in access across the country. PCGs are local groupings of general practices that are involved in the commissioning of local community and secondary services and may eventually become Primary Care NHS Trusts, providing community services and commissioning secondary services. More recently still, the concept has been developed of "Care Trusts which could provide primary care and some secondary services, such as mental health services". The quality framework involves a three-layer approach¹⁹. Clear standards of service will be set by National Service Frameworks and a National Institute for Clinical Excellence (NICE), which will evaluate new treatments. An SF for Older People was published in 2001²⁰. The first of eight standards was "rooting out age discrimination". The seventh concerned mental health and included NICE guidelines for anti-dementia drugs. The new NSF made it clear that standards in the Mental Health NSF already published applied to older people. Local delivery of services will be made dependable by a combination of lifelong learning²¹ linked to professional self-regulation and clinical governance. Clinical governance²² places obligations on Chief Executives of NHS Trusts to make arrangements to monitor and continuously improve the quality of health care they provide. Finally, all this will be underpinned by the national monitoring of standards involving a National Performance Framework, an inspectorate (the Commission for Health Improvement) and a National Patient and User Survey. This ambitious vision sets a massive agenda for change and demands radical shifts in the management and clinical cultures of the NHS of a magnitude that will not easily be achieved²³. Without adequate resources, these well-intentioned reforms may well overload the capacity for change of both managers and clinicians working in the NHS.

Primary care groups (in England) result in a much greater influence for general practitioners and other primary care workers in the commissioning of secondary services. Their boards are dominated by primary care workers. When they become Primary Care Trusts, good corporate governance demands a board structure with executive and non-executive directors, more analogous to the boards of existing Hospital and Community Trusts. This will result in general practitioners effectively losing their quasi-independent status. The Trusts are likely to be direct providers of many community services currently provided by Community or Community and Mental Health Trusts. In Scotland, where different arrangements pertain, mental health is part of primary care groups, but they do not control the budgets of secondary care Trusts in the same way as in England. In Wales the arrangements are closer to those in England. In England and Wales it seems likely that, in the larger cities at least, "stand-alone" Mental Health Trusts will be the order of the day. For old age psychiatry, this probably means that the managerial separation between old age psychiatry and geriatric medicine will be perpetuated. Unless imaginative and pragmatic solutions are found, this could result in many demarcation problems. However, a new culture of collaboration rather than competition and a government apparently committed to encouraging "joined-up" thinking and working means that these difficulties may be overcome.

The funding of the NHS is probably even more important than its organization in determining the future of health care in the UK. It was the squeeze on NHS development in the 1980s that provoked the medical profession to campaign for more development money. In the light of international comparisons, both of spending on health care and the age structure of the population, this campaign seemed fully justified. The previous reorganization increased management costs and reduced the morale of many in the NHS. So far, although the new government has promised more capital investment in the NHS, it has continued to support the controversial Private Finance Initiative (PFI) as a main strand of funding, which reduces the Public Sector Borrowing Requirement. There is concern that PFI will result in a reduction in bed numbers and diversion of money away from services to support repayments to private providers in respect of capital developments. If more money for services comes with the new reorganization, then the potential for positive change exists. If it does not, then the new reforms, like the Internal Market before them, are doomed to failure. Another problem with introducing improvements is the shortage of trained staff. The government has recognized the need to train more doctors and nurses, but it will take time to turn this recognition into staff-delivering services.

OTHER INNOVATIONS

The direct reforms of the NHS have been accompanied by a Royal Commission to review long-term care. The report of this Commission²⁴ controversially suggested that the personal care and residential elements of continuing care should be separately funded. Personal care should be paid for from general taxation, whilst living and housing expenses should continue to be means-tested and subject to co-payment. The government has yet to make an unequivocal positive response to this. Standards in psychiatric services for old people continue to improve generally but it remains to be seen whether the latest reforms will ensure that quality is universally high and that funding is adequate.

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The Pattern of Psychogeriatric Services

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HISTORIC BACKGROUND

The roots of the National Health Service (NHS) and the development of psychogeriatric services in the UK are discussed in the previous chapter. The evolution of psychogeriatric services has been guided by professional knowledge and opinion, by the politics of the health and social services, by financial constraints and, occasionally, by public opinion. From the inception of the NHS, which effectively antedated the beginning of provision of specialist psychogeriatric services until the 1990s, there was a consensus about how developments in services should occur in response to changing demography and epidemiology as well as advances in medical knowledge.

This consensus was threatened in the UK by the imposition of the ideology of “market forces”. Much long-stay hospital accommodation was effectively “privatized” by decisions to support patients in private nursing and residential homes from state funds and to close down as many long-stay NHS beds as possible. The ideology of market forces was briefly applied to local authority provision of community care. Since April 1991, psychiatric patients with social needs have been subject to a “care-planning procedure” in which all parties, including social services, have to agree. In April 1993 the full implementation of the Community Care Act made local social services departments responsible for purchasing continuing nursing and residential home care, largely from the private sector and with a limited budget.

In the late 1990s, a government came to power that did not share the vision of market forces as the best way to regulate the NHS. This government emphasized *equity* and *quality* and returned to a modified vision of the NHS as a centrally regulated nationalized service industry. It is not certain that the opposition shares this vision, and so there is a danger of the NHS remaining a “political football”. For the time being, though, despite the distractions provided by problems in under-resourced psychiatric services for working-age adults, it seems likely that the conditions in the NHS will again be more favourable to the growth of old age psychiatry services, which are essentially a collaborative rather than a competitive enterprise.

THEORETICAL BASIS FOR PSYCHOGERIATRIC SERVICES

The pioneers of specialist psychiatric services for old people were motivated by the increasing need for psychiatric services for the age group, consequent upon increased life expectancy,

the growing knowledge base about psychiatric disorders amongst old people, and the success of geriatric medicine. The special needs of older people were not always recognized by the generic services. Diagnostic problems included the differential diagnosis of dementia, the association of apparent cognitive impairment with some cases of depressive illness, and the non-specific presentation of disease in old people. The multiple pathology suffered by old people led to a need for new patterns of multidisciplinary working and for close liaison with physicians in geriatric medicine and social services¹⁻³. As in the early days of geriatric medicine, assessment and treatment in the community were emphasized not only because of “blocked beds” but also because a more realistic picture of the patient’s health problems usually emerged. More recently, advances in psychosocial care⁴, interest in the spiritual needs of old people⁵ and the advent of new classes of antidepressant, antipsychotic and antidementia drugs (discussed elsewhere in this volume), have all had their impact on the organization and delivery of psychiatric services.

CARE OR TREATMENT—PRIMARY OR SECONDARY?

One of the key theoretical issues for the future development of community services is likely to be the distinction between care and treatment. “Care” is a word with many connotations. Some are positive but, in the medical world at least, some are negative. For example, “care” is seen as what is provided when there is no possibility of effective treatment, as in the “prescription” of “tender loving care” for the terminally ill person. “Care” tends to be relegated to untrained (although not necessarily unskilled) workers employed by Social Services, whereas “treatment” is the province of highly trained personnel employed by the Health Service. The move to “Care in the Community” may serve to reclassify older mentally ill people as not needing medical treatment, and this will have to be resisted vigorously.

This situation is further complicated by the tendency of some health planners to equate primary care with *low cost and community care*, and secondary care with *high cost and hospital care*. Old age psychiatry services straddle the hospital–community divide and provide essentially secondary services, largely in a community setting. The new term, “intermediate care”, describes well some of these community services, but some who use the term believe that community psychiatric nursing services should be part of “primary care”, when in fact they work most effectively as part of secondary community care.

KEY COMPONENTS OF PSYCHOGERIATRIC SERVICES

Catchment Area and Comprehensiveness

Virtually all psychogeriatric services in the UK work to a defined geographical catchment area and the vast majority aim to provide a comprehensive psychiatric service to all people over the age of 65 years^{6,7}. Many services are now also trying to provide for people with early-onset dementia, although often without any dedicated resources⁷.

The Multidisciplinary Team

For some this is an outmoded concept, for others an ideal that cannot be obtained, but for many psychogeriatricians it is an essential context for all their endeavours. Most multidisciplinary teams for the elderly incorporate *community nurses*, a *social worker*, one or more *occupational therapists*, a *physiotherapist*, and often a *psychologist*. Various patterns of working have evolved and been described but they have in common an attempt to involve all disciplines in formulating treatment plans for the patient.

Home Assessment

This lies at the heart of most psychogeriatric services. Surveys^{6,7} have shown that around two-thirds of referrals were seen at home by a doctor, one-fifth by other members of the team and one-tenth in outpatient clinics. Just over one in 10 were seen as liaison referrals, although in some services this rises to a quarter or even one-third, perhaps partly depending on the admission policies of local geriatric services. Less than one in 20 were admitted direct without prior assessment.

Community Treatment

The rate of acute admissions was only one-third of the rate of referrals, reflecting the fact that most home assessments do not result in admission but in treatment in the community. Home visits by community nurses are probably the commonest form of treatment in the community, although home visits by doctors and other members of the multidisciplinary team also play an important part.

Day Hospitals

In 1985, there were about 1.2 day hospital places/1000 elderly people, and this had not changed significantly by the mid-1990s. Some services and Health Regions had relatively more and others less. The use of day hospitals varied from area to area, depending on the resource availability locally. Anecdotal evidence suggests that government guidelines overestimate the need for dementia places. In many but not all cases of dementia, the need is for *care* rather than *treatment* and so a proportion of this day provision can be provided by Social Services or voluntary agencies. Here, however, issues will have to be addressed as to what kind of care is of most benefit to older people with dementia and their relatives, neighbours and friends. Elderly people with functional illness often have problems with psychiatric or psychological management that demand the *treatment* resources of a true day hospital.

Acute Inpatient Beds

The national rate of provision in 1985 was around 1/1000 elderly served, and again did not vary much over the next 10 years. This may be insufficient to cope with the increasing demands caused by demographic changes and the relative loss of long-stay beds but a great deal depends on the community services available, since there is potential for considerable "marginal shift" between community and inpatient resources. One study showed that around a quarter of acute psychiatric beds for all age groups were occupied by elderly people with depressive illness, and in many areas anecdotal evidence suggests that a greater proportion of acute psychiatric beds are being used for functional illness, principally depression. As with day hospital places, it appears that the old guidelines may have overestimated the needs for dementia assessment beds and underestimated the needs for patients with depressive illness and other functional illnesses. Because of the high prevalence of physical illnesses in mentally ill old people, it is recommended that acute beds should be on a general hospital site. Some services are now beginning to differentiate the assessment and management of behavioural problems in demented people, which can be carried out in the community or in community-based units, from the management of patients with depression, who often have major associated physical illness or disability (and may need ECT) and are therefore better managed on an acute hospital site. The same may apply to atypical dementia patients requiring high levels of investigation or to patients with dementia and delirium. This last group may be best helped on geriatric medical wards.

Long-stay Beds

The provision in 1985 was around 3.4 beds/1000 elderly. Since then a large number of beds appear to have been closed, with patients discharged to the private sector, where developments have been funded through the Social Security budget. In 1996 the number had reduced to around 1.1 beds/1000 elderly. Since provision is largely (but not exclusively) for those with severe dementia, whose main need is for care rather than treatment, this development appears to demand a cautious welcome. Patients are generally being cared for in smaller units. However, there must be reservations. The smaller units are harder to inspect and they are not necessarily in the patients' communities of origin, since planning permission and housing costs enter into the commercial equation. They are subject to capricious changes in the market, including government refusal to pay the "going rate". They are not under specialist medical management and there is some evidence that this management may be one of the factors that reduces the rate of decline in demented elderly people, a factor which, if confirmed, might also have relevance in the day care setting. The switch to private care has been engineered for political reasons and its impact is yet to be fully assessed. Psychogeriatric services will need to retain a proportion of their long-stay beds for rehabilitation, for treatment of old people with resistant functional illnesses (especially depression) and for treatment of behavioural disturbances amongst demented people. A survey of old age psychiatrists⁸ showed the majority in favour of around 1.5 long-stay (including respite) beds/1000 elderly in community NHS units, with national rather than local eligibility criteria. The use of such beds for respite care to support carers in the community is now well established, and there is a potential for developing community units as centres of excellence for dementia care, as well as bases for multidisciplinary community teams.

GUIDELINES

The first guidelines for provision of psychiatric services for old people came in a Department of Health circular in 1972⁹. The government White Paper, *Better Services for the Mentally Ill*¹⁰, in 1975, suggested that services for old people should often be provided by a psychiatrist with “a special interest” and incorporated guidelines for bed and day hospital provision for the “elderly severely mentally infirm”. Subsequently, the Royal College of Psychiatrists produced guidelines intended to help College representatives reviewing job descriptions for new or replacement consultants. These were endorsed by the Health Advisory Service in its report, *The Rising Tide*¹¹, and by the joint Royal College of Physicians and Psychiatrists report, *Care of Elderly People with Mental Illness*¹². The second joint report¹³ adopted a more multidisciplinary approach and described the different types of mental illness in old age as well as appealing for equity and “national reference frameworks”, which corresponded closely to the concept of “National Service Frameworks” introduced by the new government¹⁴. It cited “indicative service levels”.

REALITY AND GUIDELINES

Although in the UK, old age psychiatry is now recognized as a specialty by the Department of Health, we are still awaiting the collection of routine statistics. Initial manpower statistics appear to be grossly inaccurate. The most comprehensive data available are from the 1985 survey, updated by the 1996 survey, which unfortunately did not achieve such wide coverage. There are also now agreed international standards for old age psychiatry services¹⁵ and an international survey has established that basic levels of service exist in 12 countries worldwide¹⁶.

SPECIALIST SERVICES OR NOT?

In view of the documented rapid expansion of services over the years, this question may seem superfluous. However, it was possible as a result of the 1985 survey to compare services where psychogeriatricians work half-time or more in the specialty with those where the consultant commitment to the elderly is less than half-time¹⁷. Specialist services had generally higher staffing ratios (with the exception of non-consultant medical staff), a higher proportion of acute beds on general hospital sites and a greater proportion of long-stay beds within the area served. These last two could be regarded as surrogate indicators of quality of care. In addition, the specialist psychiatrists were more likely to look after all mental illness in old age—the recommended pattern of service—to engage in teaching and to show an interest in research.

CONCLUSION

Psychogeriatrics has “come of age” in the UK. Provision, although geographically patchy and relatively under-resourced, still provides one model for future developments. This model, fostered in the National Health Service with its principles of equality of access, payment from general taxation and central planning, survived the major changes of the 1990 “reforms” and should thrive under the regime of equity and quality proposed by the present UK government, always provided that old age psychiatrists show adequate leadership and that governments furnish adequate resources.

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Organization of Services for the Elderly with Mental Disorders

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In 1997, the Section of Geriatric Psychiatry (now Section of Old Age Psychiatry), World Psychiatric Association, with the participation of the Division of Mental Health and Prevention of Substance Abuse, World Health Organization, met at Lausanne to develop a Consensus Statement¹ on the *Organization of Care in Psychiatry of the Elderly*. Under the very able leadership of the late Jean Wertheimer of Lausanne, a multidisciplinary group of leaders in the field produced such a statement. This chapter briefly abstracts and highlights the central issues contained in this document with my personal annotations and explanations in italics.

GENERAL PRINCIPLES

Any organization of care must be founded in firm principles of human rights of the elderly with mental disorders. The historical discrimination towards the mentally ill should be actively fought and the destigmatization of mental illness must be conducted in the context of the unalienable human rights of each individual.

- Good health and life of good quality are fundamental human rights. This applies equally to people of all age groups and to people with mental disorders.
- All people have the right of access to a range of services that can respond to their health and social needs. These needs should be met appropriately for the cultural setting and in accordance with scientific knowledge and ethical requirements.
- Governments have a responsibility to improve and maintain the general and mental health of older people and to support their families and carers by the provision of health and social measures adapted to the specific needs of the local community.
- Older people with mental health problems and their families and carers have the right to participate individually and collectively in the planning and implementation of their health care.
- Services should be designed for the promotion of mental health in old age as well as for the assessment, diagnosis and management of the full range of mental disorders and disabilities encountered by older people.
- Governments need to recognize the crucial role of non-governmental agencies and work in partnership with them.
- Preparing for increasing life expectancy and ensuing health risks calls for significant social innovations at the individual and societal level, which must be founded on a knowledge base drawn from contributions by, and collaboration among, the medical, behavioural, psychological, biological and social sciences.
- In developing countries it may be difficult to provide resources for the provision of care. This, however, does not invalidate the aims of helping the elderly by the application of the principles listed above and the specific principles that follow.

SPECIFIC PRINCIPLES

The acronym CARITAS was deliberately chosen as a cogent reminder that "care and love" underpin these principles. The word "systemic" in some service contexts may be replaced by the word "seamless", representing a service organization that does not permit each individual to fall between gaps.

Good quality care for older people with mental health problems is: Comprehensive, Accessible, Responsive, Individualized, Trans-disciplinary, Accountable, and Systemic (CARITAS).

- A *comprehensive* service should take into account all aspects of the patient's physical, psychological and social needs and wishes and be patient-centred.
- An *accessible* service is user-friendly and readily available, minimizing the geographical, cultural, financial, political and linguistic obstacles to obtaining care.
- A *responsive* service is one that listens to and understands the problems brought to its attention and acts promptly and appropriately.
- An *individualized* service focuses on each person with a mental health problem in his/her family and community context. The planning of care must be tailored for, and acceptable to, the individual and the family, and should aim wherever possible to maintain and support the person within his/her home environment.
- A *trans-disciplinary* approach goes beyond traditional professional boundaries to optimize the contributions of people with a range of personal and professional skills. Such an approach also facilitates collaboration with voluntary and other agencies to provide a comprehensive range of community-orientated services.
- An *accountable* service is one that accepts responsibility for assuring the quality of the service it delivers and monitors this in partnership with patients and their families. Such a service must be ethically and culturally sensitive.
- A *systemic* approach flexibly integrates all available services to ensure continuity of care and coordinates all levels of service providers, including local, provincial and national governments and community organizations.

COMPONENTS OF SERVICE

The concept of "surround with care" as represented by Figure 1, of concentric cycles, provides the concept of service delivery to have as its core the patients and their family carers. By enveloping them with seamless service organization with "permeable" boundaries represented by "dotted lines" in the figure, they are totally supported, and with changing circumstances can move from one service component to another without hindrance.

The described components of service, A–H, when put into place effectively, will meet the international best practice expected by this Consensus Statement.

The components of services can be summarized in Figure 1, which portrays the concept that individual patients, together with family and carers, are surrounded by the care service; these are flexibly interlocking, overlapping and integrated to provide a unified system for continuing care and best possible quality of life. Structural obstacles are minimized, as represented by the dotted lines of the figure, enabling the smooth movement of the patient from one service component to another as changing circumstances require. This section describes the components that can be put into place to address the care needs described in the previous section.

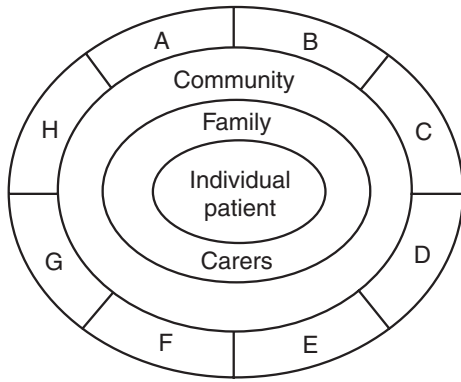


Figure 1 Surround with care

The following components ideally should be the responsibility of specialized teams of trained healthcare professionals working in psychiatry of the elderly. Where there is a scarcity of trained staff and resources, it will be necessary to use *ad hoc* solutions in order to provide the necessary components—while trying to develop services fully.

1. *Community Mental Health Teams (CMHTs) for Older People.* The lead in organizing the following components of the service should ideally be taken by multidisciplinary specialist teams working on psychiatry of the elderly. The CMHT may consist of doctors, psychiatric nurses, psychologists, social workers, therapists and secretaries. Referral to the CMHT is usually from primary care. One of the main responsibilities of the CMHT is the specialist assessment, investigation and the treatment of people in their home setting. In situations where such personnel are not available, the responsibility may be taken by general psychiatric or geriatric medicine teams.
2. *Inpatient services.* Acute inpatient units need to provide specialist assessment and treatment for the full range of mental disorders. This may in some cases include rehabilitation before return to the community.
3. *Day hospitals.* This is an acute service which offers assessment and treatment to older people who can be maintained at home, supported by the multidisciplinary team. The day hospital team could include doctors, nurses and therapy staff. Transport may need to be available.
4. *Outpatient services.* These provide assessment, diagnosis and treatment for people fit enough to live in the community and get to and from the hospital base. Outpatient services should be close to the inpatient and day-patient units. They may involve subspecialty clinics (e.g. memory or mood disorder clinics) and mobile clinics.
5. *Hospital respite care.* Hospital beds may be used to provide a respite service for people with chronic and severe mental illness and associated difficult behavioural problems, in order to give their carers a break and enable care at home to continue as long as possible.
6. *Continuing hospital care.* Care for life in a hospital setting may be required for people with chronic and severe mental illness and associated difficult behavioural problems. Such care should be provided in as relaxed and homely an environment as possible, with carers encouraged to participate.
7. *Liaison services.* Consultations and/or liaison services should be provided between facilities for elderly people with mental disorders and those serving general and geriatric medicine, general psychiatry, residential facilities and social agencies. This relationship should be of a reciprocal nature.

8. *Primary care.* The primary care team has the initial responsibility for identifying, assessing and managing mental health problems in older people. The decision to refer to the CMHT is usually made in primary care.
9. *Community and social support services.* These are services (both formal and informal) to enable the elderly person to remain at home, including a range of activities (home care, day care, residential care, respite care, self-help groups, etc.) provided by voluntary or government/social services.
 - (a) *Respite facilities.* A range of short-term, time-limited, in-the-home and out-of-the-home services (residential services, other carers, day programmes) to support the carers.
 - (b) *Residential care.* For those patients whose physical, psychological and/or social dependencies make living at home no longer possible, a spectrum of residential facilities should be provided. These range from supported accommodation with low-level supervision, medium-level care facilities, to full nursing facilities. These should be organized to achieve the best possible quality of life.
10. *Prevention.* The mental health team for the elderly should engage in the prevention of relapse of disorders by careful follow-up. They should also identify the risk factors for mental disorders in the elderly (e.g. hypertension, alcohol and substance abuse) and ensure these are effectively managed by appropriate medical, social strategies. Within each service, preventive activities need to be coordinated in collaboration with relevant public health and other healthcare professionals. These may include educational activities to improve early identification of mental health problems by carers, families and primary care personnel in the community.

CONCLUSION

While it may be seen to be less than realistic for some economically less-advantaged countries to achieve this best-practice ideal, nevertheless, both the WPA and WHO would recommend that such an ideal be both aspirational and inspirational. The paragraph:

In developing countries it may be difficult to provide resources for the provision of care. This, however, does not invalidate the aims of helping the elderly by the application of the principles listed above and the specific principles that follow

in the General Principles section is reiterated here as a reminder that all available resources should be deployed within limited resources. For the “developed” countries that fail to meet these ideals, the WPA and WHO strongly urge their governments, policy makers and healthcare professionals to strive towards satisfying and exceeding the described principles and components of service.

As development of services is always a dynamic process, it is hoped that, in the future, this consensus statement will be reviewed and extended to incorporate improvements and innovations.

The central philosophy is that all elderly people with mental disorders should be provided with the best quality of life and the best quality of care in recognition of their esteemed status as Elders in our society.

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The Multidisciplinary Team

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Teamwork in the animal kingdom is a behaviour that enables survival. One example of its success is the male emperor penguin, who incubates his precious single egg balanced on his feet for 60 days through the total darkness of the Antarctic winter. This deed is achieved by the emperors huddling together and shuffling around in a constant movement, providing ever-changing relief from the gale-force wind¹.

The ability to behave in such a team has evolved through generations of practice. In care of the elderly in human populations there has been no such tradition and, for those suffering from psychiatric illness, the need for effective teamwork was never more urgent. The problem list presented by the functionally or organically ill elderly patient is often disconcertingly long and complex. The psychiatric disorder cannot be isolated from physical health problems, neither can the patient's functioning be assessed outside the context of his/her own domestic and social situation. Thus, in attempting to treat the patient it is necessary to combine the skills of psychiatric and geriatric medicine, together with those of nursing, clinical psychology, remedial therapy and social work. In many situations, input from the Housing Department, voluntary agencies and the local chaplaincy will be required. The creation of community care packages defined by the UK Government White Paper, *Caring for People in the Next Decade and Beyond*, highlighted the development of teamwork in case management². The UK Government has extended this strategy to include carers as partners of professionals in the planning and delivery of care³. Teamwork in the management of the detained patient, particularly in respect of discharge planning and aftercare, are recommended as good practice by the Department of Health⁴.

No one profession can attempt to provide all the various skills necessary in the management of every case. It is the relationship between the different professions that is so important. The concept of the multidisciplinary team can be described as a group of members of different professions whose working skills, when combined for the needs of the patient, aim to exceed in quality the simple summation of their individual abilities. Teamwork adds that extra vital ingredient, which needs further exploration.

The multidisciplinary team should be restricted in size to those personnel actively involved in management of cases and in close geographical proximity, to enable face-to-face contact between workers. However, the team should not exist in isolation from other services. The services that collaborate most successfully internally have been shown to not necessarily work best with other agencies⁵. Between different agencies there should be formal liaison and agreed policies over borderline cases and service demarcations.

FEATURES OF SUCCESSFUL MULTIDISCIPLINARY TEAMWORK

Communication

The Personal Social Service Council, in its review of community care, found that failures in communication and negotiation with other agencies led to ignorance of the roles and skills of other professional groups. The organization of care was less good in the more complex situations. As a result, wrong courses of action were taken by some professionals, including inappropriate admissions to both hospital and residential care. The need for joint planning was stressed, particularly between professionals of the different agencies at the operational level⁶. Effective communication is grafted onto the rootstock of mutual trust and respect for the individual roles and skills of other team members. Autonomous decisions, unnecessary duplication of work and stereotyping of one professional by another are thus avoided.

It is important that communication within the multidisciplinary team is afforded formal expression in regular meetings. Too much "corridor" decision making can lead to mistrust. Regular attendance by all personnel complements the identity and strength of the team. This results in the creation of a suitable arena for the airing of grievances and resolving of disputes. Teamwork involves allowing the patient and carer to participate in the process. The therapeutic team could include the patient⁷, although the current climate of consumerism in care might be perceived as threatening by some workers.

Leadership

Traditionally, the leadership role in health care is assumed by the senior doctor; this in part reflects society's expectations and faith in the medical profession. Treatment directives may be instigated by the initial medical assessment and form the basis for the leadership role in the clinical team. The dominant role of the doctor in the healthcare team may have a stabilizing effect on the team's structure and prevent leadership struggles by other members⁸. Too hierarchical a structure, though, will result in too many decisions made by one person. More democratic teams will develop more flexible work practices and collective decision making. The doctor may in fact be the least qualified in management and leadership skills within the team. Moreover, the reason why multidisciplinary teams can work so badly is frequently the scant regard paid to them by the medical profession. Non-medical leadership of the multidisciplinary team can work, particularly if the issue of responsibility for

decisions is satisfactorily addressed. A major DHSS document is clear that a consultant may not be held responsible for negligence on the part of others, simply because he is the "Responsible Medical Officer". It states he is not accountable like a military commander; "the multidisciplinary team has no commander in this sense"⁹. There is, however, a tendency to equate responsibility with out-of-hours accessibility, especially during a crisis.

It is important to remember that the key worker may not be the same person responsible for providing alternative care in the event of a crisis; e.g. the community mental health nurse (CMHN) may be the link with the elderly person at risk, but is unable to access directly residential care facilities should they be needed urgently.

For whoever dons the mantle of leader, skills in communication are a vital requirement, as is the ability to create a feeling of mutual trust and respect in order to maximize members' strengths and create compromise between individual members to fit in with overall team goals. Much of the motivation for the team will depend on the qualities of leadership. The creation of realistic goals gives a sense of purpose and a framework. Obviously, resource constraints are a major barrier to this process. It is important to distinguish the demands of budget holding from clinical management, and where the multidisciplinary leader is also the budget holder, the team members will need to learn the wider issues in the face of restrictions caused by tight fiscal control. They will need to feel the leader's commitment and sincerity in the care of their clients or patients.

Audit

The Department of Health encouraged medical audit as a means of improving service delivery and management¹⁰. Care of the elderly psychiatrically ill requires more complex assessment than pure medical audit, and the multidisciplinary team is in the key position to instigate clinical audit because of its potential for knocking down interprofessional barriers and prejudice. Multidisciplinary audit can be attempted if the process is viewed from a positive and non-defensive position. Good and bad practices in liaison will be easily revealed, particularly if the patients' and carers' views are included in outcome measures. The team should be in a constant state of evolution and be able to incorporate new ideas. This sense of objectivity and self-assessment will be created by a positive clinical audit programme and will itself become part of the quality of the successful team.

Morale

Clearly, morale is dependent on the key issues—leadership, communication, achievement of goals and self-assessment, described above. Other ingredients need adding to the recipe. Morale relies heavily on personal support and encouragement within the team. Through time, most team members experience stresses and problems, which may affect their judgement and possibly their self-esteem. This support may be available discreetly or offered more formally through staff groups and supervision. Whatever its form, its value is its ready accessibility and confidentiality.

The ability of the team members to see the funny side of a situation is often a vital component in the maintenance of morale. Involvement in non-clinical activities, such as fundraising or games matches against other units, may have positive effects on morale—even in defeat. In essence, it is important to create in team members a sense of personal value and ownership of their team. In the current climate of health service reorganization and cost-benefit assessment, the sharing of successes in patient treatment can contribute greatly to good morale.

FUTURE DEVELOPMENT OF THE MULTIDISCIPLINARY APPROACH

The introduction of goal-orientated multidisciplinary methods of treatment require changes in structures of healthcare organization¹¹. Sadly, the move towards care management detailed in *Caring for People in the Next Decade and Beyond*² has not been complemented by sufficient funding to meet the requisite planning, training and research needs. Either the resources are not available to get multidisciplinary teams properly established, or there is insufficient funding to achieve even modest treatment goals. It is disputes over limited funding that can lead to the break-up of collaboration between agencies and effective teamwork.

Planning

The last decade has, however, seen an encouraging trend in the sophistication of expertise within the psychogeriatric team, producing greater quality of care of the elderly psychiatrically ill. The CMHN has developed roles in early case assessment and liaison with associated professionals. Because of the high profile in the locality, the CMHN has forged links with many agencies, e.g. district nurses, specialist housing, residential care facilities, and facilities for the treatment of alcoholism. Together with the increasingly skilled work with dementia sufferers, functionally ill patients and carers, the CMHN now brings to the multidisciplinary team a more vital and complex role. The social worker is still not given sufficient time to use her training in casework from the all too demanding accommodation-finding role. Historically, the champion of patients' rights, the social worker, may now access independent advocacy schemes. The roles of the occupational therapist and physiotherapist have been further strengthened by their involvement in community assessment liaison. The clinical psychologist finds much demand within the team, not only for assessment of areas of cognitive deficit in the patient, but also increasingly for the treatment of functional disorder, e.g. cognitive therapy and the evaluation of staff management methods and attitudes. There remains a paucity of training places in clinical psychology.

The shifting nature of the multidisciplinary team approach is further advanced by the recognition of the increasing role the voluntary sector can offer, both informally, using volunteers as befrienders of the patient, and more formally by such organizations as Age Concern and the Alzheimer's Disease Society, providing essential services such as day care. The role of the coordinator of such services can play an important part within the multidisciplinary team¹².

The expanding therapeutic team must recognize the problems that may arise when a new worker comes to fill a specific role carried out more generally by other workers, e.g. a social worker being appointed where previously much informal casework was conducted by the CMHN. Members of smaller teams may appear to have less distinct professional roles. Concentration on individual strengths, not weaknesses, will enable expansion of the team to take place.

Training

It is not clear that such growth in individual professional expertise in care of the elderly psychiatrically ill has been matched by the requisite training in multidisciplinary teamwork skills, particularly in undergraduate medical education. The General Medical Council, in its publication, *Teaching Tomorrow's Doctors*, has recognized this shortfall¹³. Attitudes and techniques in

cooperation with other disciplines will need to be explored in order to escape from the purely medical model of treatment.

Case studies have been described as being well suited for teaching, in an interesting and understandable way, the complexities in the field of geriatric medicine, where an increased degree of integration with other services is required¹⁴. This must also be true in geriatric psychiatry. By the careful selection of cases to illustrate psychological, medical and social issues, the trainee should be stimulated rather than confused. This is an ideal opportunity for multidisciplinary teaching, enabling the learner to understand the different professional roles. The aim should be to show that complex cases might be manageable by the establishment of treatment goals, even if they are necessarily limited. Cultural, ethical and legal issues can be shown to have an important influence on the equation. Teaching by the use of case studies can be a good focus for multidisciplinary learning for groups of trainees from different professional backgrounds.

Planned community visiting sessions with the CMHN or social worker are highly valued by undergraduates of all relevant professions at Southampton University Medical School.

Doctors in higher training to be consultants in the field of old age psychiatry demand experience in multidisciplinary teamwork and should have flexibility in their programmes to permit in-depth exploration of the roles of the different members of the team. A variety of multidisciplinary teams should be observed, including those in other psychogeriatric units.

Individual professional training must be mirrored by developments encompassing different professional groups. A training model has been devised to overcome the organizational isolation of agencies serving mutual elderly clients with mental health problems in Philadelphia¹⁵. Training issues were explored at a conference on interdisciplinary issues in mental health and ageing, where interprofessional cooperation was reaffirmed and strengthened¹⁶. There is a rising swell of workshops on collaborative care in the management of the elderly psychiatrically ill, much of it promoted by initiatives from the voluntary sector and much of it relevant to the training needs of the multidisciplinary team.

Research

Objective research in the workings of the multidisciplinary team has received little attention in the case of the elderly psychiatrically ill. A review of the role and effect of this teamwork in the management of delirium in dementia found insufficient data for the development of evidence-based guidelines¹⁷. A difficulty is that evaluation has been mainly restricted to descriptive terms using, for example, in-depth individual case studies. In this area in

particular, clinical audit and operational research are almost indistinguishable. Outcome measures, however, using quantity and timing of liaison, do not measure the essential components of the workings of the team—the quality of the relationships and the ability to respond as a whole to the needs of the patient. Study is still required into the methods of evaluating the effect of the quality of these relationships, particularly within the field of old age psychiatry.

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Community Care: the Background

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The specialty of old age psychiatry in Britain grew out of the failure of orthodox, institutionally-based psychiatry to meet the needs of the growing number of elderly people with dementia. Its success stemmed from its proactive community-based approach, an emphasis on the support of family and other carers and the demonstration of the positive outcomes obtainable through better recognition and energetic treatment of much of the functional illness of old age¹. Crucial to that approach and the effectiveness of the elderly mental health service itself has been the development of partnerships across health and social services, support to informal carers and the increasing range of care staff working directly with patients. Just as it has led the rest of psychiatry in assertive outreach, it has often taken over from the increasingly hospital-centred geriatric medicine as a catalyst and advocate for community care for older people. This chapter will look at some of the components and contributors to that care, as well as some of the wider organizational and political factors that have influenced it.

PRIMARY HEALTH CARE

General practice has always been the cornerstone of the National Health Service (NHS) in Britain, providing continuity of individual and family care and acting as gatekeeper to secondary care. Aggregation into group practices paved the way to the establishment of the primary healthcare team (PHCT), with its core membership of general practitioner (GP), practice nurse, district nurse and health visitor, and later a looser attachment or sessional input from community mental health nurses (CMHNs), podiatrists, physiotherapists, counsellors and specialists from secondary care. The PHCT provides the bulk of domiciliary health care and surveillance of those at risk and the main day-to-day link with social services, housing and other local agencies supporting disabled people in their homes.

With the aim of strengthening the locality and primary care focus of the NHS, the incoming Labour government² has aggregated groups of PHCTs covering “natural” localities of 50–150 000 population into primary care groups (PCGs). Healthcare budgets for these localities have been devolved to PCGs, who are responsible for increasing the equity and quality of primary care within local populations, for assessing their health needs and for commissioning secondary care services. There is also the option/expectation that PCGs will progress to fuller autonomy as primary care trusts (PCTs). These will be empowered to take over the management, for instance, of community hospitals and appropriate secondary care services and will be expected to develop closer partnerships with local social services. It is likely,

therefore, that PCTs will take on a much more comprehensive role in the provision of community care within the Health Service. Parallel legislation^{3,4} will also facilitate partnership with local authority services by enabling them to pool and ring-fence budgets to promote joint services, long advocated by proponents of community care.

INFORMAL CARERS

By far the largest contribution to domiciliary care, particularly for those with dementia, is of course that of family members and friends. The main carer is usually a spouse or partner, although care by daughters, sons and daughters-in-law remains substantial, despite greater geographical mobility, changing employment patterns and the steadily falling ratio of middle-aged to elderly people. Input from neighbours can also be vital and some areas have well-developed community networks. The House of Commons Social Services Select Committee⁵ estimated the “replacement cost” of informal carers for the disabled in Britain (over three-quarters of a million “on their feet” for more than 50 h week) as well above the total expenditure on all statutory care. The government recently acknowledged the importance of carers and the load they carry and announced a strategy to ensure that their needs will be properly assessed and that they become eligible for extra financial support⁶.

BACK-UP FROM GERIATRIC PSYCHIATRY

Partnership with family carers is fundamental to the practice of old age psychiatry, especially in the context of dementia. Initial assessment should include the extent of informal care, the health and attitude of the carers and the prospects for sustaining such care in the future, given adequate help. Carers need the support of a key worker with a good knowledge of the “system” who can help them access appropriate services and allowances and the practical and psychological value of carer support groups. Important aspects of that key working are its continuity, attention to the health and morale of the carers and prompt arrangement of further assessment, intervention and respite when the need arises.

For those with psychiatric illness, and particularly dementia, the key worker will usually be the CMHN, working closely with colleagues in primary health care and social services teams. CMHNs also have an important support and educational role, with the increasing number of untrained voluntary and paid carers working with patients in their homes, day centres and residential care. They will also draw on the range of skills and

resources of the specialist team itself for further assessment, treatment, respite or crisis intervention. A great deal of the work of the elderly mental health service takes place in the patient's home. Ours, for example (catering for an elderly population of 60 000), has 20 CMHNs, all with large case-loads and between them clocking up about 20 000 patient contacts a year. This is complemented by outreach from other members of the service (e.g. 8000 medical community contacts), two day hospitals and deployment of staff to support day centres specializing in the care of the elderly patients with mental health problems. An original bed complement of 250 (mainly long-stay) has reconfigured to 40 continuing-care and 90 short-stay beds, 35 of the latter for functional illness. This has necessitated extensive outreach and support to residential and nursing homes but has released funds to redeploy into community developments, such as the establishment⁷ of a sitting service to offer home respite to carers for demented patients. This has now diversified into other areas of community support for elderly people with mental health problems as a thriving provider in the new mixed economy of community care.

SOCIAL CARE IN THE COMMUNITY

Progress in community care in Britain has been hampered by the very different way in which health and social services are organized and financed and the frequent lack of co-terminosity in their local boundaries. The NHS is funded from taxation and is predominantly free at the point of delivery. It has strong central direction and is coordinated locally by (unelected) district health authorities. Social and housing services are run by (elected) local authorities, which receive block allocations from central government but also have to raise much of their revenue through local taxation and by means-tested charging of services to clients. Local authorities were traditionally the main providers of domiciliary and residential support for disabled people.

The 1970s saw many initiatives in specialized supported housing for older people⁸⁻¹⁰ and in individualized and sometimes intensive domiciliary services to support, in their own homes, frail elderly people who might otherwise have required residential care. Evaluation of these and subsequent projects usually showed them to be valued by clients and generally cost-effective at low and moderate levels of disability, but decreasingly so (and at times very expensive) for those with heavier dependency^{10,11}. Because they generally rely on intermittent carer input or on the client summoning assistance, they tend to be less effective for patients with advancing dementia than for those with a purely physical disability.

Generalization of these initiatives in community care was seriously hampered by the economic problems of the 1970s and early 1980s. The incoming Thatcher government found things reaching crisis point but was unwilling to give the necessary support to local authorities to expand domiciliary based care. Instead, it changed the regulations governing social security benefits¹² to allow these to be used to pay fees in private residential and nursing homes, with no provision for needs assessment. This led to a huge expansion (from £10 million to over £1 billion a year during the 1980s) in this sector, but none in domiciliary provision, which was not eligible for this source of funding. This wasteful and retrograde trend was castigated by the Audit Commission¹³, which drew attention to the contrasting effectiveness and popularity of the Kent Community Care scheme^{14,15}. This had been using care management to assess the needs of frail elderly people in the community and to put together packages of domiciliary care, drawing on a range of neighbourhood resources. This was enabling many people who would otherwise have needed residential care to remain in their own

homes, despite a budget ceiling well below the cost of the residential placement.

To establish a transition to this model, the Audit Commission proposed a ring-fencing of all public expenditure on community and institutionally-based continuing care (including all residential, nursing home and NHS continuing care) to be administered by a single district care agency on a care management basis. These proposals were endorsed by the government's management guru¹⁶, who also stressed the need to ring-fence the budgets for all continuing care and to ensure adequate transitional funding to secure ample early investment in domiciliary-based options. The resulting White Paper¹⁷ unfortunately ignored these two crucial points. The result gave Social Services departments the responsibility for care management (including the purchase of long-term nursing home care) but without arrangements for any transfer of funds released within the NHS if it closed equivalent continuing-care beds.

Implementation of the Act since 1993 has therefore brought about a modified resurgence of community care for older people in Britain. Social services departments were encouraged to focus on the assessment and care management functions and to stimulate a mixed economy of domiciliary care providers, from whom they purchase packages tailored to the needs of clients. It is very gratifying to see one's patients getting individualized support packages, combining informal, social and healthcare elements and often assembled at short notice in the event of a crisis. Aware of the growing pressure on acute hospital beds, some health and social services departments have also collaborated to develop domiciliary and intermediate care options to reduce the need for hospital admission and facilitate early discharge¹⁸. The advent of PCGs should certainly stimulate that sort of approach. Unfortunately, because the means-testing rules make residential care cheaper for social services than the domiciliary equivalent for anyone with moderate care needs, the cost ceiling applied by most departments to domiciliary packages for individual clients is a long way below the level of residential home fees. This rules out such care for many people for whom it would otherwise have continued to be cost-effective, particularly those with dementia.

POLITICAL AND ECONOMIC FACTORS

Lack of investment has been the main cause of our slow progress with community care for older people in Britain. The politicians have generally been seduced into thinking that it is a cheap option, but at times of economic hardship domiciliary provision has always been the first target for cuts. In the 1980s spending on care of the elderly increased greatly but in the wrong direction and it is taking a long time for the pendulum to swing back. The separate organization and funding of health and social care have created perverse incentives, which have been exploited to preserve an inadequately funded NHS by narrowing its focus to the care of acute illness. This constriction of the range of free NHS care has led to artificial and increasingly absurd demarcations between health and social care, which undermines efficient and effective partnership and has thrown an increasing share of the cost onto patients themselves. Failure to ring-fence the NHS continuing care within the community care budget has allowed the closure of 60 000 beds since 1990 (with savings redeployed to support other NHS priorities), while social services have had to pick up the load with an increase in their nursing home purchasing from 100 000 to 180 000 beds—a cost shift to social services and (through means testing) to patients and families of over £300 million a year. Not surprisingly, this has encroached on investment in domiciliary care, which only increased by 56% in the first 5 years of implementation of the community care legislation¹⁸.

This shrinkage in the role of the NHS in continuing care and rehabilitation and the barrier to effective services created by the artificial distinction between health and social care¹⁹ has been widely criticized^{18,20,21}. There is growing consensus that things will only get properly “joined up” when the funding rules for health and social care are harmonized and the nettle of the cost of our ageing population is properly grasped by the politicians. The Royal Commission²³ set up by the government recommended that all care (as opposed to board and lodging and general living) costs should be publicly funded from general taxation, eliciting little response as yet from a government still wedded to reducing taxation and maintaining means testing. The earlier Joseph Rowntree Enquiry²² reached similar conclusions on the care costs but favoured a hypothecated, prospective, compulsory health/social care insurance levied on payrolls as a way of enabling each generation to fund its own care in old age. If the government finally accepts one or other of these options, the millennium could usher in an exciting new phase in community care.

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Tom Arie

University of Nottingham, UK

The Nottingham University Department of Health Care of the Elderly was designed as a collaboration in which physicians, psychiatrists, gerontologists and other health workers are equal partners. It is neither a department of psychiatry with geriatricians on its staff, nor vice versa: it is an integrated joint enterprise. It was the model for 20 years. Following the retirement of the foundation professor, the university restructured the Department but the service ethos continues, as does the joint teaching programme. Described here is the Joint Department as it was, and as it became well known, from 1977 to 1997.

Diagrams may put things best. Figure 1 shows the structure, and makes the important point that although the department is unified, its services are differentiated. Physicians do medical work, psychiatrists psychiatry. There is cross-training, and above all constant formal and informal collaboration and support, both in the hospital and in assessing and keeping people going at home.

The aim is to make easily available what patients need. Thus, a patient of the psychiatric service has as easy access to physicians and their facilities as if he/she were their patient, and vice versa. In this way it is possible to offer responses that match the pattern of morbidity characteristic of the very old, namely that it is mixed and often unpredictably changeable. There are no demarcation disputes, and there is no waiting list. Things were very different when the department was established in 1977.

INTEGRATION

The services are based in two general teaching hospitals. In University Hospital the professorial unit has a medical and a psychiatric ward side-by-side. There are close links with the rest of the health specialties and professions, both inside and outside the hospital. For instance, trainees rotate from general medicine,

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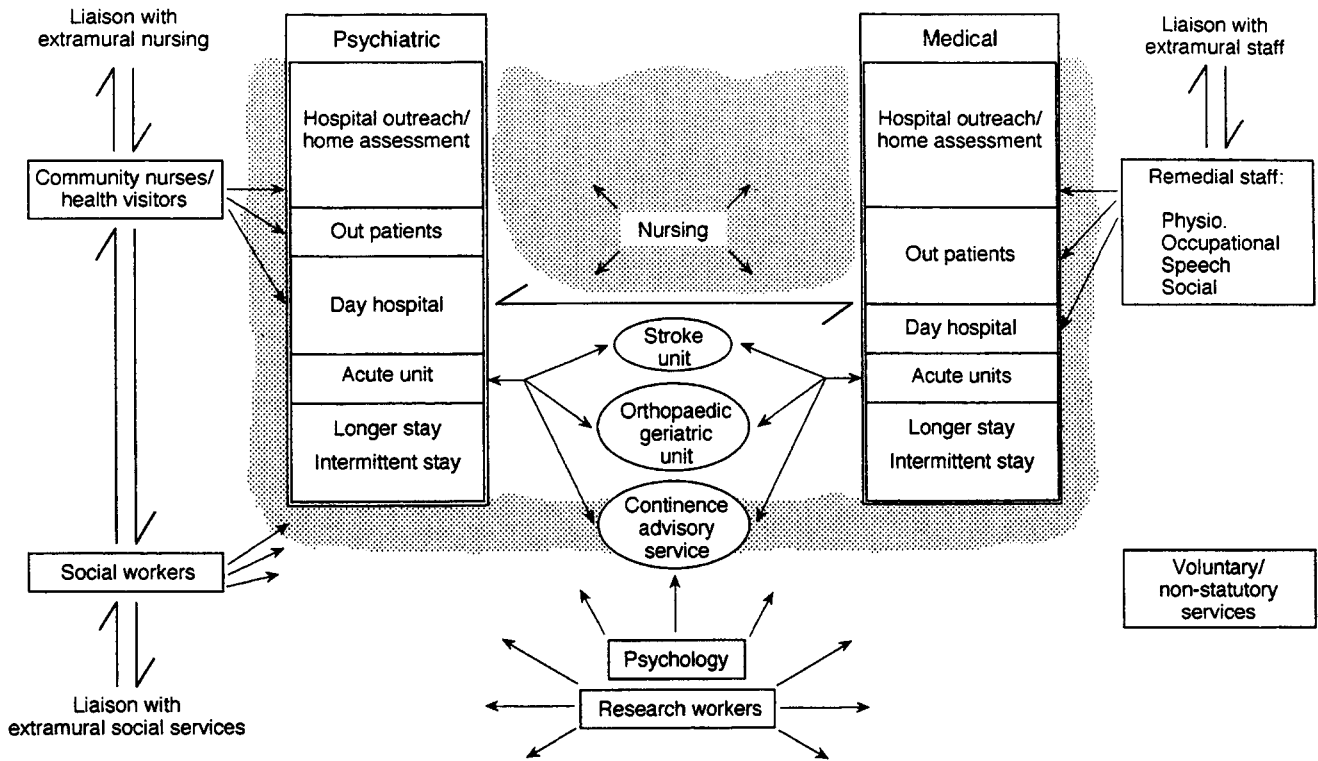


Figure 1

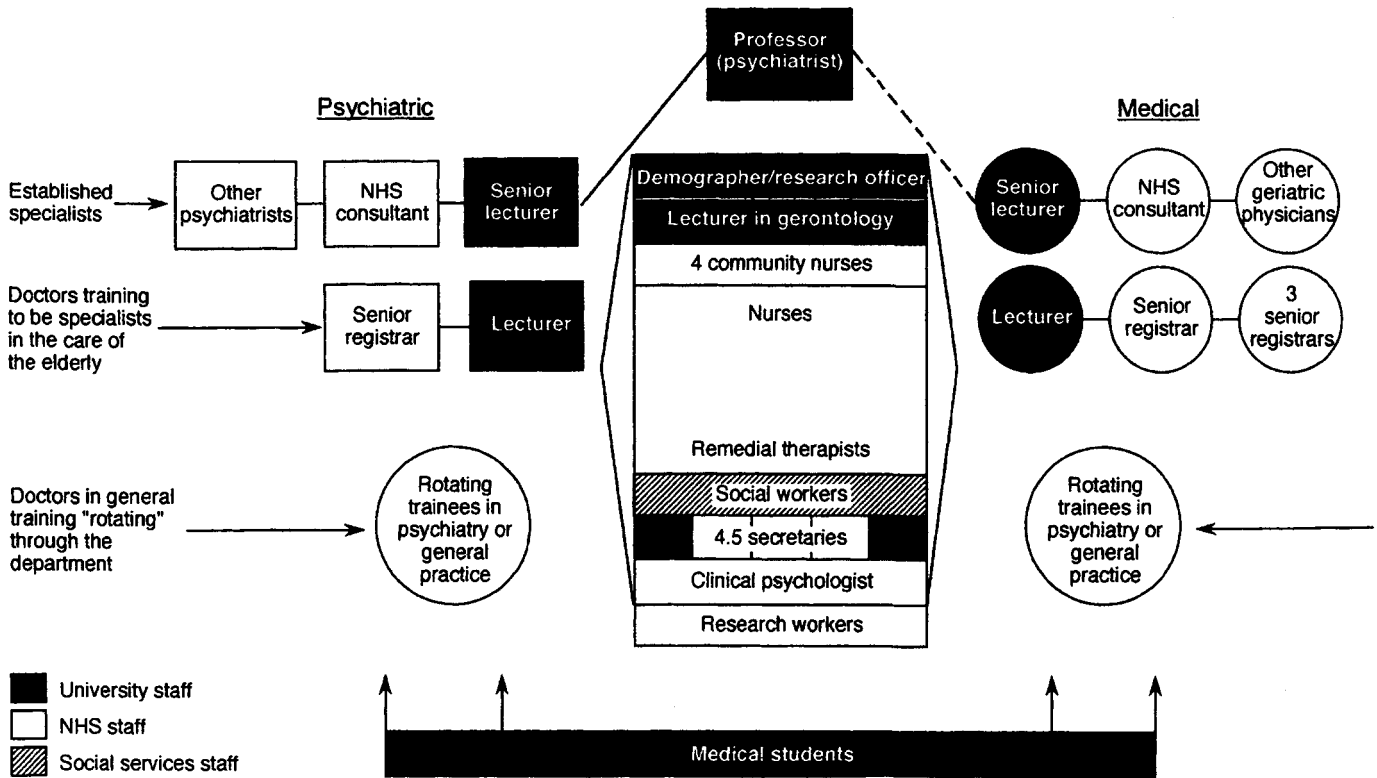


Figure 2

psychiatry and general practice (Figure 2), and there is pairing with medical and surgical firms. The department is embedded in the local medical community, and particularly values its links with the parent disciplines of medicine and psychiatry and its closeness to primary care. All referrals to the psychiatric side of the department are initially seen at home by a senior psychiatrist, usually together with a medical student, and any other team members or community staff who are involved and able to be present, and if possible the family doctor is present too. In practice, most visits are by psychiatrist and medical student. On the medical side of the department a home assessment service is freely available, but the majority of patients are seen in the clinic, or admitted as emergencies. Community nurses, working outwards from the hospital, are part of both the medical and psychiatric services.

EDUCATION AND RESEARCH

All Nottingham medical students spend a month full-time in the department. Teaching is a joint effort between all staff, of whatever specialty, medical and non-medical. The course comprises: a clinical clerkship, inside and outside the hospital; a planned course of teaching; and a personal project under supervision. With about 150 students a year, there is a heavy teaching load. The course is well received and many students come back as postgraduate trainees. Rotating trainees come to acquire better skills (and, we hope, greater job satisfaction) in dealing with old people in whatever field they decide eventually to work. Only

the senior registrars are being trained to become specialists in old age.

Research thrives on collaboration and transcends boundaries of departments: we have long been collaborating with other departments, and within a mixed department research collaboration comes easily. Major projects include longitudinal studies of well-being of old people at home—the Activity and Ageing Study—and participation in the Medical Research Council study of Cognitive Function and Ageing (CFAS). Most projects have involved staff from both of the main branches of the department.

CONCLUSION

What has been called “the Nottingham Model” works well, and is a satisfying way in which to work. There have naturally been day-to-day problems, some general, others parochial, but none that have divided the department along specialty lines. Above all, patients do not fall between stools and bucks do not get passed.

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The Development of Day Hospitals and Day Care

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ORIGINS

Day hospitals are said to be among the few notable creations that psychiatry has given to medicine¹. The concept that older people could be given treatment, rehabilitation and care without resorting to “inpatient” status within hospitals or related “institutions” was accepted enthusiastically in the UK from the late 1960s. “Partial hospitalization” (usually day hospital care, but sometimes night care, relief admission programmes, rotating care or shared bed schemes), linked to active community services, has been a feature of the services provided for the elderly, not only by psychiatrists but also geriatric physicians. Non-medical agencies, such as social services departments and voluntary bodies, have also participated. Interestingly, and perhaps significantly, the private sector has been much less enthusiastic for partial hospitalization schemes. This appears to be true for other countries.

ATTRACTIONS

The attractions of day care are most obvious for the patient, who is enabled to maintain his/her home routines and contact with his/her supports, whilst taking advantage of professional expertise in treatment during certain parts of the day. Family members or carers may have mixed perceptions: their burden of anxiety and responsibility for the ill or disabled patient is relieved only episodically and only in part; yet many are pleased with this opportunity to continue to contribute to care and for therapy with the guidance of professionals. The resources of the day hospital add to the resources of family life². For the health care agency (the National Health Service), day hospital development has been attractive for its flexibility and apparent cheapness. It may not, however, represent a cheap option. Its importance to health care systems lies in its influence on the effectiveness and smooth running of inpatient and other domiciliary or extramural and partial hospitalization projects. These are increasingly likely to be cooperative ventures with other care-providing agencies, such as social services³.

SPREAD WITHIN THE UK

At the end of the 1960s, the Department of Health was prompted by a pressing need to review the services it was providing to the increasing numbers of older people suffering from dementia and other major psychiatric disorders. This revealed a woeful situation, with relatively few beds, badly supported and in

inappropriate locations, frequently operating with waiting lists, that saw patients dying before admission⁴.

A review of provision based on the 1971 census was to establish the baseline for inpatient services⁵ and health authorities were encouraged to create day hospital places in equal numbers, often starting from zero⁶. As little guidance was given in respect of the development of these day hospitals, a wide variety of interpretations arose from practitioners and planners trying to produce the best possible facilities. Between 1976–1986 the total number of day hospital places for the mentally ill increased by 50%. Provision for the elderly ranged from 129 places/10 000 in East Anglia to 693/10 000 in the North West⁷. Wide variations persist between areas in respect of day hospital places available to the elderly populations served, and the Royal College of Psychiatry’s recommendations of one place for every 500 (or fewer) of the elderly population⁸ is not yet seen everywhere, for a variety of financial and other reasons.

STRUCTURES

The cheapest option for developing day hospital provision was to place the facility in a disused part of a mental hospital, and risk the deficiencies of the parent hospital transferring to the new unit. Interestingly, the effect tended to be positive, for both the new venture and the existing facilities for this patient group, and illustrates a potential of such care—the link with inpatient services benefits the day care unit, its attenders and their carers, and encourages wards to provide care more relevant to population needs. Alternatively, developments took advantage of existing premises, designed for other purposes but away from the mental hospital and often in the area to be served⁹. “Travelling” day hospitals used a network of such premises, staff moving between the locations to provide care on different days¹⁰, a useful system for rural communities.

The ideal would be purpose-built units, probably on health authority land and sited conveniently in the catchment area. No bed provision is required and staff are not employed when the unit is closed. These developments can also create a positive image for a vital, if unfashionable, service by raising its community profile.

Such “stand-alone” day hospitals run the risk of being seen as distant from the inpatient services, and may detract from the reputation of the latter, especially as they are often seen as more thrusting and challenging. This can hinder cooperative efforts, such as when a patient requires a period of short-term, rotating or prolonged inpatient care¹¹. Probably the best model places day hospital and inpatient facilities within the same unit, with several

such units of modest size at strategic sites throughout the community to be served (Figure 124.1).

FUNCTIONS

A day hospital may aim to provide one or more of a series of care options for the elderly attender. It may offer assessment and treatment otherwise requiring admission, facilitate early discharge from inpatient care, provide rehabilitation, or help attenders deskilled or disabled by long experience of illness. Additionally, it may provide continuing support for patients with established disabilities (with or without a known vulnerability to major decompensations), arguably in itself a rehabilitative regimen attempting to prevent further deterioration and also providing carer support. In practice, many day hospitals perform mixed functions and may not have formally determined aims. The experience of others can serve to inform the process of defining such objectives.

Bergmann *et al.*'s¹² general hospital-based day hospital in Newcastle functioned as an assessment and early treatment unit, without initial home visits and very much in the manner of a tertiary referral facility for other teams working out of existing mental hospitals. Alongside was a small ward with a similar

function, and the team also undertook liaison work—probably representing the best use for this day hospital within the city's then mental health provision. Even so, and despite a highly skilled and single-minded team, the successful pursuit of “assessment only” is difficult to sustain. Bergmann reported that the day hospital's advice, after assessment, was taken or acted upon in disappointingly few cases. Whitehead¹³ also stressed the assessment and treatment potential of the day hospital, designating it perhaps the key element in a community-orientated old age service. He was, however, providing a comprehensive range of services within one team, with home-based assessment and treatment available before a decision regarding the use of the day hospital or a small inpatient unit. His unsatisfactory experience of poor-quality mental hospital wards determined his philosophy that recourse to inpatient care would be taken for very few patients.

Such models of care are attractive, but were influenced by other factors: for Bergmann, longer-term care was available elsewhere, and Whitehead had a supportive nursing home sector sympathetic to his vision¹⁴. Other authors have been uncomfortable with day hospitals that appear to function mostly to delay entry into long-term care^{15,16}; however, for some individuals and their carers, this may be the very help they want, and should be evaluated in this light.

Arie¹⁷ identified two groups of patients benefiting from long-term day hospital attendance: dementia sufferers living with carers wanting them to remain at home, but who needed help and some respite from the carer role; and patients with relapsing illnesses complicated by persisting “neurotic” symptoms when well. The latter were almost all women living alone. Others working in well-sited and well-equipped day hospitals have confirmed these images of the successful long-term patient. Here, the male:female ratio of dementia sufferers was often equal, and some attended from rest homes. There was also an intermediate group, suffering from mood disorder or neurotic symptoms associated with quite severe physical disability (from stroke, parkinsonism, arthritis, chest or heart conditions), having nursing needs as great as those of dementia sufferers, but with psychological needs akin to other “functional” patients¹¹.

The “mix” of day hospital attenders can be crucial. Individuals severely damaged by dementia benefit from simple, structured and repetitive activities that are unhelpful and unrewarding to their cognitively intact peers. Where dementia has caused a decline in physical abilities, basic group physiotherapy techniques can be beneficial, whereas other patients with complex physical and psychological needs may only thrive with individually tailored personal care interventions. Mixing patients with widely differing cognitive abilities can be unacceptable to both the better and the less able. Potential tensions can be reduced by using different areas of the unit for different purposes, or by having day-specific activities. Where several local day hospitals coexist, they may develop specialized roles¹¹.

Most practitioners with day hospitals available for their patients are enthusiastic about their potential, but there have been few robust evaluations of the efficacy of day hospitals. Philpott's review¹⁸ cautiously concluded that day hospitals remained experimental care delivery systems. Creed and colleagues, after reviewing studies of day hospital care¹⁹, evaluated day hospital activities in the north-west of England²⁰. They confirmed that day hospitals are able to provide alternative care for many patients but that not all such facilities can, or want to, provide the same service²¹. Longer periods of treatment may be offset by greater efficacy in preventing relapse and maintaining independence²².

The costs of day hospital care are probably lower than those for inpatient care. MacDonald *et al.*²³ undertook an extensive study of outcomes in matched groups of elderly people with mild to moderate dementia in four care settings in London. They found no significant differences in mortality or in changes on dementia scores. They concluded that non-hospital day centres offered the best value.

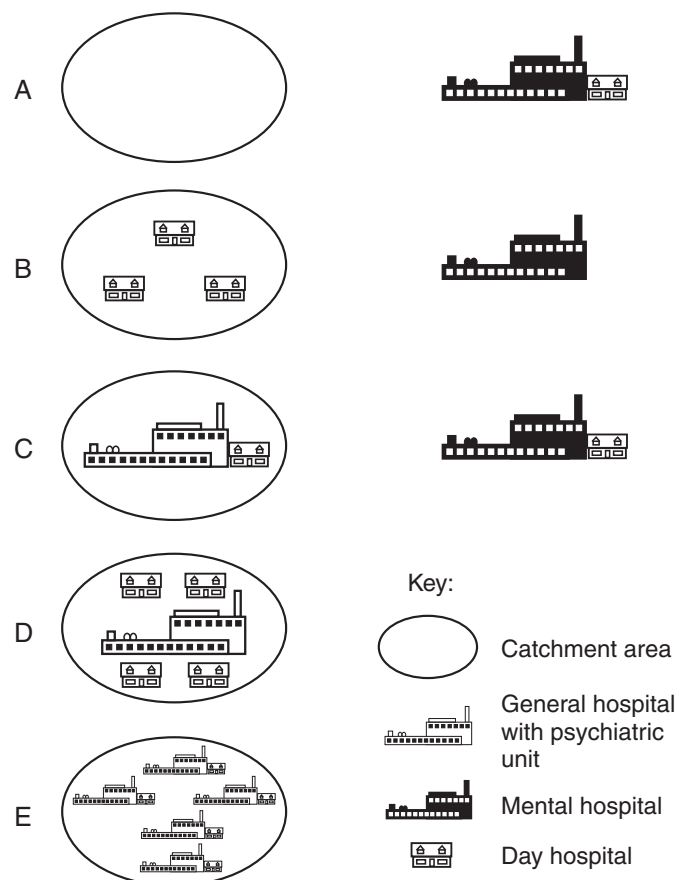


Figure 124.1 Models of day hospital provision. (A) Large mental hospital geographically distant from catchment area served, starting day care on site. (B) Mental hospital supported by stand-alone day hospitals within catchment area. (C) Mental hospital beginning to give way to district hospital unit, both sites with inpatient and day hospital facilities. (D) General hospital with psychiatric unit supported by stand-alone day hospitals. (E) General hospital with day hospital unit and supported by several units within the catchment area, each offering inpatient and day hospital care.

Panella *et al.*²⁴ found that only 69 of 314 elderly demented patients likely to benefit from day hospital programmes in Burke, USA, took advantage of them. For these, the costs were estimated to be considerably lower than nursing home care and the programme was acceptable to patients and carers. Zeeli and Isaacs²⁵, despite describing a geriatric day hospital where 75% of attenders' time was spent in active therapy and transport problems were non-existent, still showed only half the patients completing planned treatment. Only one-third achieved staff-determined objectives or felt improved by 3 months.

Transport problems, along with changes in health or social circumstances, were often associated with costly lost days of care in the comprehensive review performed by Eagle *et al.*²⁶, but the work confirmed the positive effects of day hospital care on the physical and emotional health of elderly attenders.

The Audit Commission²⁷ has confirmed the expense of day care when linked to domiciliary support services to the moderately disabled—it can exceed the cost of residential care, and the cost of informal care is all but impossible to calculate. This does not mean it is not preferable to other forms of care.

Donaldson, Wright and Maynard²⁸ emphasized the need to consider several measures when assessing day hospital care. These included the costs of all services used over a designated period, with clinical, social and psychological measures at set intervals related to treatment and follow-up, and carer and patient satisfaction scores. As yet, few if any studies have examined all these aspects of care, although a work in progress by Read and her colleagues in a joint day care/day hospital project in Wolverhampton is designed along very similar lines to those suggested by Donaldson's team²⁹. This evaluation is also attempting to place such care in context with other care and services available. More comprehensive assessments of the efficacy of all available complementary facilities individually and within systems are necessary to tease out the particular effects of day hospitals. These would include measures of the morale and self-image of professional carers in the sectors and their views and attitudes towards other complementary services, patients and their carers. These considerations reflect the complexities of the processes of care delivery to the elderly mentally ill, and the relationships between "consumers" and "providers".

CONCLUSION

Day hospitals for the elderly mentally ill have become established services, especially within the UK. There is little reason to doubt that they fulfil a useful and acceptable function. Providing something of this nature has both energized and encouraged developments in many areas. Refinements of practice occur over time as individual day hospitals modify what they do in the context of a changing awareness of their own potentials and weaknesses, and in relation to changes in other services available to the potential client or patient group. Formal research evaluations continue and may better inform such evolutionary developments and future planning. It remains important to be aware of the limitations of such evaluative research and it would be quite wrong to stop good ideas being put into practice on the "quasi-scientific" basis that "its cost-effectiveness cannot be demonstrated".

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Day Care

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The first psychiatric day hospitals in the UK opened in the late 1940s and their numbers have continued to grow over the five subsequent decades. In geriatric psychiatry, as in geriatric medicine, the proliferation of day care facilities came rather later, with large expansions of such services since the 1960s. A similar proliferation of day care facilities for the elderly with psychiatric difficulties has occurred in other parts of the world¹⁻³, with parallel expansions in both the health and social care sectors⁴. Since their inception, day care facilities, in all their many and disparate forms, have come to be viewed as essential components of most comprehensive old age psychiatry services. This chapter will seek to follow a brief description of day care facilities with a discussion of their aims and effectiveness, concluding with some current issues and possible future directions for day care services.

MODELS OF DAY CARE

Models of day care for the elderly mentally ill are extremely diverse. Indeed, given the relative lack of critical evaluation of such services, this is perhaps to be expected. This diversity is enhanced by the range of bodies funding and running day care services, including local authorities, the health service, voluntary agencies and the private sector.

Day hospitals and day centres are often described as distinctly differing facilities (as they will be in this chapter for the sake of clarity), although in reality the two types of facility more probably form part of a continuum. Day hospitals tend to be funded solely through health service budgets and to be staffed by health service professionals. Day hospitals will often be linked geographically with a hospital site and will thus have access to a full range of physical treatment and investigative facilities. The programmes at day hospitals are often intensive⁵, with reality orientation, formalized social interaction, occupational therapy, physical activities and frequent meetings with the patients' families. Day centres, by contrast, are funded more usually by local authorities and/or voluntary agencies, and have a much lower proportion of health service professionals on their staff. Day centres are optimally sited in the community, close to the patient's home. The programmes at day centres tend to be less intensive and "medicalized", with greater emphasis on informal socialization and diversionary activities. Almost self-evidently, day centres are significantly cheaper to run than are day hospitals.

There is debate as to the optimal "mix" of patients/clients in day care facilities. Most day hospitals do mix the demented and the functionally ill elderly, although the latter group would usually form only 10–20% of the total. Since the first day

hospitals were established, the numbers of functionally ill patients have probably declined, not through explicit policy decisions but as a result of increasing pressure from the growing number of demented elderly patients⁶. Some regions tend to incorporate the functionally ill elderly into general adult day hospitals, feeling that this affords such patients an environment that can be more accurately tailored to their needs. Chodosh *et al.*⁷ describe a day care programme that mixes physically infirm with demented patients, but these authors reflect that their mentally infirm attenders would probably be better suited to a separate programme, in view of their shortened attention spans. In areas of low population density, taking services out to the patients, through travelling day hospitals, can be a very useful innovation^{8,9}. In more rural areas, especially since underutilization of expensive custom-built facilities can have an adverse influence upon cost-effectiveness¹⁰, it may be appropriate to integrate demented patients requiring day care into small community-based inpatient units¹¹.

AIMS OF DAY CARE

While it is fair to say that there is a general lack of clarity and agreement as to the aims of day care for the elderly, certain themes would be common to most patients in most settings. These would include relief from loneliness and boredom and the provision of stimulation and social support¹², while the aims of day hospitals will usually be broader and more ambitious.

As noted above, the substantial majority of elderly day hospital attenders suffer primarily from dementing illness. Most day hospitals, therefore, are geared more towards the needs of the demented than of the functionally ill elderly, whose day care requirements have thus received rather less attention. Cited roles of day hospitals for the functionally ill include assessment, continuation of treatment after hospital care, monitoring and encouraging compliance with medication, and focused group activities dealing with issues such as bereavement.

For the demented elderly, many aims of day hospital care have been described and these would include: assessment, providing an alternative to institutional care, treatment of cognitive and non-cognitive symptoms of dementia, relieving stress on caregivers, determining whether institutional care is required and what the optimal form thereof will comprise, easing the transition from home to institutional care, shortening hospital stays, acting as centres of expertise from which skills can be "exported", enhancing the morale of old age staff who rotate through day hospitals, and providing a focus for liaison with other care providers. The issue of whether these aims are effectively fulfilled will be discussed below.

It may sometimes be the case that the role of a day care facility is more reactive than proactive, and one pragmatic view is that the shape of day care develops partly to plug gaps in local services¹³. It may be worth noting also that the literature about the aims and benefits of day care is written almost exclusively by health care professionals about patients; certainly among younger patient groups, the most valued aspects of day care may not accord at all closely with those that might be highlighted by staff¹⁴.

EFFECTIVENESS OF DAY CARE

There has been no evaluative research on the effectiveness of day care for the functionally ill elderly. If one can validly extrapolate from findings in younger patients, then compliance with medication can be enhanced by focused interventions with day hospital attenders¹⁵. Similarly, extending findings from the demented elderly might suggest that the often very stressful activity of caring for a depressed or psychotic relative would be relieved by the patient entering day care.

While it remains an under-researched area, there has been more evaluation of the effects of day care attendance among the demented elderly, although many of the hypothesized functions of day care have not been meaningfully scrutinized. For example, while many old age services regard assessment as a pivotal role of their day hospitals^{9,16,17}, another view is that this can be conducted just as well as an outpatient and/or in the patient's home¹⁰. There are no data to support either standpoint.

Reports of specific cognitive or non-cognitive benefits of day care among demented patients have not been consistent or confirmed, and tend to be based on anecdote¹⁸. Where ratings of change are made by carers, then such ratings may be influenced by improvements in the carers' well-being as a result of their dependants' day care attendance. If day care is indeed specifically beneficial to patients, then the improvements may be more subtle than can be measured with existing rating measures, and Donaghy¹⁹ has noted that a validated quality of life measure for the demented elderly would constitute a significant advance. It is perhaps noteworthy that professionals now working with the demented elderly predict that day care would enhance their own quality of life should they develop dementia²⁰.

While day hospitals developed largely as an alleged alternative to long-term inpatient or other institutional care,¹⁷ this hypothesized role may be outdated and there is little evidence that they fulfil this function. Patients admitted into long-stay care following day hospital attendance have been found to be no more disabled, cognitively or behaviourally, than those who had no experience of day care²¹, suggesting that there was no delay in institutionalization. New day care facilities in an area do not appear to change the requirements for long-stay care¹⁶. It may be, however, that day care can shorten stays in acute inpatient facilities by acting as something of a "half-way house" between hospital and home while community links are re-established¹⁷. Mintzer *et al.*²² in South Carolina, found that there was equal improvement in the agitation of demented patients who had a short admission followed by home assessment plus "partial hospitalization" and those who experienced a lengthier hospital stay.

The strongest evidence for the effectiveness of day care is in improving the well-being of carers. Earlier studies, such as that by Gilleard²³, tended to be naturalistic but demonstrated improvements in the psychological well-being of carers after their dependant commenced day care attendance. These improvements were not related to changes in the problems presented by their dependant and seemed to derive, therefore, from "time out" from the stresses of the care-giving role. More recently, quite well-designed controlled studies have been conducted. In Australia, Wells *et al.*² conducted a "before and after" day care comparison

of supporters, and also a comparison of supporters receiving and those waiting for day care. They found no differences in caregivers' well-being but reflected that the average period of relief (11.9 hours/week) may have been insufficient to confer benefits. In a recent large study, Zarit *et al.*²⁴ found that supporters of demented day care attenders (minimum of 2 days/week) felt less overloaded, strained, depressed and angry than did supporters whose dependant had no access to day care. The issue of what the "effective dose" of day care might be to enhance carers' well-being remains a fairly open question¹⁸.

COST-EFFECTIVENESS OF DAY CARE

Given the relative dearth of evaluative studies on the effectiveness of day care, it is unsurprising that cost-effectiveness is also an understudied area. With regard to the dementing elderly, the Scottish Chief Scientist Working Party²⁵ concluded, in 1988, that "the available research indicates that virtually nothing is known about the economics of day hospitals", and since then little useful information has accrued¹⁸. The study by Mintzer *et al.*²² mentioned above, did conclude that their use of "partial hospitalization" was clearly a cost-effective method of instigating and continuing treatment for agitated patients with dementia. In Sweden, Wimo *et al.*³ compared 55 demented patients in day care with 45 waiting list controls and noted a statistically non-significant trend suggesting that day care may be more cost-effective than other types of service provision. In their review of 16 adult day centres across the USA, Reifler *et al.*²⁶ considered that one of the keys to cost efficiency was improved utilization, which related to meeting "customer demands", such as the need for transport and opening for a longer day between 7.30 a.m. and 6 p.m. Perhaps specific to health and social care systems in the USA, these authors make other suggestions as to how the financial viability of day care facilities can be enhanced.

DAY HOSPITALS OR DAY CENTRES?

Studies of demented attenders at day hospitals and at day centres tend to find that those at the former are rather more demented and more behaviourally disturbed, but that the similarities are more striking than the differences between the two groups of attenders²⁷⁻²⁹. This gives rise to the view that a proportion of demented attenders at day hospitals could have their needs met more cost-effectively at day centres, as has also been suggested for younger adults attending day hospitals in north-west England³⁰. This view is strengthened by findings that supporters of the demented elderly attending day hospitals do not derive more psychological benefit than those whose dependant goes to a day centre^{28,31,32}. Comparisons against day centres of other suggested functions of day hospitals, such as assessment and treatment of non-cognitive symptoms, have not been conducted. Furthermore, no randomized trials of day hospital vs. day centre care have been conducted among demented subjects, and while such studies would be difficult to conduct, they are likely to be feasible, given that they are possible among the physically disabled elderly³³.

PROBLEMS AND ISSUES IN DAY CARE

As has been indicated above, there is a considerable need to address the issue of the effectiveness and the cost-effectiveness of different day care programmes and of the specific components of these programmes, even although this is an undeniably complex task. Other common issues in day care will be touched upon below.

Within busy day care settings, it is often difficult to ensure continuity of care for those attending. This potential problem can be ameliorated by the appointment for each client of a "key worker", whose role is to coordinate the programme for that individual. This process can be facilitated by a "problem list", with a corresponding list of proposed action for each problem area. The key worker can also play a crucial role in helping to coordinate other community services: Peach and Pathy³⁴ have shown that day hospital attendance can be coupled with a reduction in other social supports without deleterious effects upon the patient. The key worker can also maintain links with other members of the multidisciplinary team who are involved in the patient's care.

Levin *et al.*³¹ have highlighted the underprovision of day care facilities, with a minority of demented individuals in the community having access to such facilities. While selection of demented patients for day care seems somewhat arbitrary²⁸, it might be considered reasonable to prioritize day hospital places for those with complex needs, where assessment, problem identification and management may lead to improvement in the patient or allow for easier caregiving. Day hospital assessment may be especially helpful in the assessment of patients who live alone, where adequate information cannot be gathered through home or outpatient assessment. In the absence of complex needs and where the main aims are stimulation for the patient and respite for the caregiver, then lower-cost day centre care may suffice. Linked to this issue is the question of how many days per week a patient should attend. Research is required into whether day care should seek to provide a lot of respite for a few selected carers or a little respite for many more carers.

The vast majority of day care programmes operate during normal working hours. It is likely that supporters would be even more appreciative of time for themselves in the evenings and at weekends^{31,35}.

SUMMARY AND FUTURE DIRECTIONS

Few mental health practitioners would dispute that "day care, whatever its source, is a key component of comprehensive services to elderly persons and their families"³¹. However, in this increasingly cost-conscious age, such an assertion will not ensure continuation and expansion of day care services without adequate evaluation of their efficacy and cost-effectiveness. Day care services will probably become progressively aligned with other community resources for the elderly and it will surely be helpful if there is increased integration of health service day hospitals and social services day centres. Unless benefit can be demonstrated from the intensive involvement of mental health professionals, particularly nursing staff, then it is likely that there will be a progressive move toward staffing day care programmes predominantly with (cheaper) untrained personnel. In addition to studies of efficacy, there is a need to investigate further the selection of patients for day care, the most appropriate hours for programmes to operate and the optimal degree of respite that should be offered to supporters of the demented elderly.

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New Technology and the Care of Cognitively Impaired Older People

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Recent years have seen a rapid growth in research and development of new technologies to improve services and enhance the independence and quality of life of older people. The actual and potential role of new technologies has been recognized by the UK Royal Commission on Long-term Care¹, by the European Commission in their COST-A5 initiative on “Ageing and Technology” and TIDE R&D programme on “Telematics for Disabled and Elderly People”. A wide range of products is being developed²⁻⁴, using new technologies within robotics, telecommunications and information processing. In this chapter, telecare concepts and applications are discussed in terms of their potential benefits for cognitively impaired older people.

TELECARE CONCEPTS AND APPLICATIONS

“Telemedicine” is now well established worldwide and refers to the use of electronic information and communication technologies to link medical practitioners and patients⁵. This could include applications to support a wide range of medical practices, such as diagnostics, patient monitoring, therapy, rehabilitation and health education. “Telecare” is a more recent concept^{6,7} and refers to the use of new technology to deliver and facilitate health and social care support services in the community.

New technology is likely to play a significant role in the community support of older people for several reasons⁸. Increasing numbers of older people have forced health and social services to look at more cost-effective approaches to care and support. Health and social care policy argues that most older people want to stay in their own homes and the improved services and enhanced levels of safety and security afforded by telecare systems may have the potential to help very disabled people to do so. Manufacturers and service providers have recognized the market potential for meeting the demand from older people and their carers for new products and related services. A number of telecare applications are available or being developed within several application areas.

Health and Social Care Information Systems

These are used to provide healthcare professionals and clients with health information. These can include simple operator-based systems, interactive websites and self-navigating information services using terminals in surgeries⁶.

Client Support

Telecare can include telephone-based services to sophisticated medical systems utilizing state of the art telecommunications. These can be used to provide emergency response, counselling, training, information and client and carer support^{9,10}.

Client Records and Care Planning

Huston¹¹ argues that medical records are a weak link within telemedicine and telecare and outlines a model for a comprehensive medical records system. The EU-funded ITHACA project aims to provide a standard IT-based system for the assessment and planning of client care.

Assessment

Home-based technology can provide patient data that can be used for assessment and the specification of treatment and care. Doughty and Costa¹² outline a system to assess the ability of elderly people to live alone in the community, using sensors to provide electronic measures of activities that indicate functional performance after discharge from hospital.

Teleconsultation

This refers to the use of technology to facilitate communication between a healthcare professional and client. For example, Whitten *et al.*¹³ report the use of a cable television interactive video system to deliver home health services from “telenursing cockpits”.

Patient Monitoring

The most widespread technologies are community alarms or personal response systems¹⁴, which allow a person to raise an alarm at a central control facility by pressing a button on his/her telephone or on a pendant worn around the neck. Recent work^{15,16} has been aimed at developing a second generation of systems that use sensors in the home and artificial intelligence to automatically detect emergencies, even when a person is unable to raise an alarm him/herself.

Smart Housing

Smart housing helps frail or disabled people to live independently in their own homes by making the home environment more manageable. Remote control by dwellers, carers or care professionals and “intelligence” built into home systems means that some of the tasks of everyday living can be performed automatically. These include systems for controlling room temperature, home security, cookers, curtains, windows and so on.

TELECARE AND DEMENTIA

Despite considerable research and development activity in all the above areas, very little research has specifically focused on the needs of older people with cognitive impairments, and this remains a neglected area^{17,18}. For example, only two of 32 projects, funded under the final phase of TIDE³, made a reference to this client group, while none of the 127 R&D projects within the European Commission’s Health Telematics programme⁵ specifically mentioned people with cognitive impairments. This situation perhaps reflects assumptions made by engineers and designers about the abilities of cognitively impaired people to use and benefit from technologies. However, there may be considerable scope for new technologies for this client group within a number of key areas¹⁹.

Supervision and Surveillance

A major concern in supporting demented older people at home is the potential safety risk involved, and the use of automatic alarms to highlight dangerous situations has received increasing attention from technologists and academics^{18,20}. For example, wandering behaviour may pose a significant risk for an individual, as he/she may get lost or be unable to cope with the potential dangers of the outside world. However, restraint may be an undesirable or unethical option for wanderers. Simple sensors can be installed to determine whether a person has left a room or building and send a message to a carer or service provider. “Tagging” devices may be useful in locating people outside the immediate home environment.

Environmental Control

The home environment can be a dangerous place for cognitively impaired people. For example, they may leave cookers and heaters unattended, while gas and electricity are potentially lethal if misused. “Smart housing” technologies may be particularly useful for people with dementia, automatically shutting off devices or allowing remote control by carers or service providers. Smart housing could also use sensors linked to computers to control household appliances, devices, lighting and heating, depending on the movements and activities of the cognitively impaired person.

Quality of Life Care

The telecare applications discussed above generally focus on safety and security. However, there may be opportunities for more positive uses of technology to enhance the lives of people with dementia. For example, using technology to identify when a person is restless and bored and then initiating a familiar or enjoyable activity, such as playing familiar music or videos. Aromatherapy rooms could emit relaxing aromas to alleviate stress, while video telecommunications could help to reduce social isolation.

Carer Support

Carer support is a key aspect of care in the community. Using telecommunications to carry out more routine tasks, such as shopping or going to the bank or post office, may provide them with more free time, relieving the “burden” of care. Access to information advice and counselling is also important.

Reminder Devices

New technology could also provide “reminder devices” to support independent living. For instance, the TIDE-funded TASC project³ aims to provide cognitively impaired people with suitable decision support software to help in carrying out household activities, social participation, communication and vocational tasks.

ETHICAL ISSUES

The discussion so far has highlighted the benefits of telecare. However, Sixsmith¹⁶ argues that new technology can be a double-edged sword and that a number of potential problems need to be addressed:

- Will telecare increase the range of care options or will it just limit choice in a different way?
- Will the technology be intrusive, leading to an inevitable loss of privacy and dignity?
- Will the use of telecommunications mean the loss of human contact in the mental health care of older people?
- Will the introduction of new technology simply reflect marketing strategies, rather than the real needs of older people?
- Will financial savings made through the use of technology be reinvested in products and services or would it be another cost-cutting exercise?
- Will new technology just be used as another form of restraint to control the lives and activities of people with dementia?

CONCLUSION

This chapter has highlighted a number of potential areas for the development and application of new technology to support the independent living of cognitively impaired older people. However, it is clear that considerable research and development work is required if ideas are to become a reality. Moreover, there needs to be a shift in emphasis within research and development to reflect the specific needs of cognitively impaired people as well as those people with physical impairments (e.g. mobility, sensory, motor, control and manipulation). For physically impaired people, the underlying basis for technological development and design is to remove the environmental barriers that turn a person’s impairment into a disability. In contrast, cognitively impaired people may have impaired abilities to understand their environment, formulate plans, carry out actions, communicate or remember what they have done or where they are. This has a number of implications for the development and implementation of telecare for this client group.

First, the inability to use a technology may be a serious limitation. Researchers and product designers will need to develop innovative ways of allowing cognitively impaired people to interact with the range of telecare systems, e.g. voice and visual interfaces, rather than traditional interfaces (keypads, etc.).

Second, the use of “smart” technologies may be a particularly useful approach for cognitively impaired people who are unable to

articulate their needs and desires. The idea of smart or “proactive” technologies involves the use of sensors and artificial intelligence to interpret a person’s observable/physical behaviours and to anticipate his/her needs from these. For example, agitated behaviour patterns may indicate that a person is hungry or wishes to go to the toilet. The telecare system could then initiate some appropriate human or machine response, e.g. send a message to a carer or provide behavioural cues to the person him/herself. The information generated by these kinds of systems has the potential to provide a much more flexible, person-centred care regimen.

Finally, it is important that the implementation and application of new technologies are grounded in a thorough understanding of the individual within his/her social and care networks¹⁷. The needs and abilities of the person may vary considerably. People in the early stages of dementia may be able to cope reasonably well with only limited care, while others may require considerable help and support. Important individual factors to consider are level and stage of cognitive impairment, ability to carry out the activities of daily living, and problematic and emotional factors, such as anxiety and tendency to “wander”. The informal and formal care networks will also determine the kinds of technological interventions that are needed. Again, this will vary from individual to individual, depending on factors such as availability of informal carer, physical proximity, carer network and capacity to provide care. The application of new technology needs to be tailored in order to complement the person’s abilities and capacities and to provide help and support to care providers.

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The United States System of Care

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OVERVIEW

In the first edition of this book Fogel characterized the US system of health care for older adults with late-life mental disorders as a “non-system of care”, plagued by irrational incentives and multiple access barriers¹. This description highlighted a system that encouraged entrepreneurial activities among practitioners and health systems, was best at delivering specialized high-quality hospital-based care, and was constrained by perverse funding mechanisms that incentivized hospital-based and institutional care (nursing homes) but disincentivized outpatient and home-based care. While the individual elements of a comprehensive continuum of care could be found, services were described as fragmented, inadequate and poorly financed. As we move from one century into the next, considerable attention has been focused on the rapid changes occurring in health care and the changes that will need to occur in order to accommodate an aging population.

Has mental health care kept pace with these developments? Has much changed since Fogel’s original assessment? Since the original report, the USA has struggled with the need to slow the growth of rising medical expenditures. The efforts to reduce global medical expenditures have centered on the application of managed care principles and practices and government cutbacks. The emphasis has been on cost reduction, not system’s integration. In this chapter we will provide an update and overview of the current system of mental health care for older persons in the USA, with a specific emphasis on the organization and financing of services. The following components and trends will be addressed: (a) The structure and organization of mental health services for the elderly; (b) fee-for-service financing of mental health services for the elderly; (c) mental health managed care for older adults; and (d) emerging and future trends in integrated services and financing.

THE STRUCTURE AND ORGANIZATION OF MENTAL HEALTH SERVICES FOR THE ELDERLY

In an ideal system, the organization, financing and delivery of mental health services to the elderly would be a seamless continuum involving acute and continuing services across inpatient, outpatient and long-term care service settings, and networked with the general medical sector. Since mental disorders are a leading risk factor for institutionalization, improving the provision of mental health services in community settings is a major focus of public policy². The following section summarizes current mental health service settings and highlights several

demonstration projects designed to provide integrated care for the frail elderly.

Psychiatric Service Settings

Primary Care

Initial access to care for older adults is usually through the primary care sector, especially for older adults without a history of severe and persistent mental illness (SPMI). Many older persons prefer to receive treatment in primary care, and this service sector offers the advantages of proximity, affordability and coordination of medical and psychiatric co-morbidity³. Many older adults may present to primary care physicians (PCPs) with symptoms of mental distress, difficult to classify in current psychiatric classification systems⁴. The most prevalent disorders in the primary care sector are depression, anxiety, anxiety symptoms, dementia syndromes and misuse of prescription medications and alcohol use. Adequate detection, treatment and referral to the specialty mental health sector remain problematic, and have been attributed to such issues as: stigma; low priority of mental health issues in patients with serious medical disorders; inadequate referral resources; complexity of patient needs (psychiatric, medical and social); and lack of time and expertise in dealing with psychiatric problems⁵. Additionally, PCPs tend to approach psychiatric disorders using a medical model, which encourages over-reliance on medications for common disorders⁶.

Despite the problems identified, PCPs will continue to be important mental health service providers. Thus, training in geriatric psychiatry and research that focuses on how to improve PCP effectiveness remains an important challenge for the coming decades^{4,7}. PCPs will also be important mental health care providers for geriatric minority populations with late-life mental disorders in coming years⁸.

Outpatient Psychiatric Service Settings

The proportion of psychiatrists reporting large geriatric case loads has steadily increased over the last two decades^{9,10}, a trend paralleled by other mental health providers, such as psychologists and social workers. More services are being delivered, in large part due to consumer demand, better treatments, an increased recognition of how untreated late-life mental illness contributes to excess disability, and more favorable Medicare reimbursement policies.

Through the American Psychiatric Association's Practice Research Network (PRN), we are just now able to describe a nationally representative profile of patient characteristics and treatments received from psychiatrists¹¹. Similar practice-based research for other mental health disciplines has yet to be launched, and only recently have data been reported that describes the demographic and treatment characteristics of patients aged 65+ treated by psychiatrists¹². Generally speaking, PRN psychiatrists provide a full array of diagnostic and treatment services for older patients. PRN provides descriptive baseline data. About 51% of patients aged 65+ in the PRN have a primary diagnosis of an affective disorder, followed by cognitive disorders (20%) and schizophrenia (19%). Older adults have more medical comorbidities and lower initial global assessment of functioning scores. About 49% of older patients were seen in outpatient settings, followed by hospital settings (32%) and nursing home settings (16%). Over 50% of patients receive both pharmacotherapy and psychotherapy, and 40% of older patients receive pharmacotherapy alone. Only 2% receive psychotherapy alone. Of patients receiving medications, over 60% of older patients received antidepressants, 40% antipsychotics and 48% benzodiazepine medications.

Data are lacking on how well psychiatrists or other mental health clinicians employ best practices or adhere to existing treatment guidelines for older adults. Future research will be needed to answer these types of questions and establish the effectiveness of treatments for subpopulations of elderly patients, such as minorities, those living in different environments and those with significant medical-psychiatric co-morbidity.

Community Mental Health Centers

Older patients with severe and persistent mental illness (SPMI) pose significant challenges for the US system of care. These individuals have long-term care needs, have limited financial resources, and secular trends in managed care, home- and community-based alternatives to institutional care are being promoted as the major venue for mental health services¹³. About 2% of persons aged 55+ in the USA have SPMI, which is expected to double over the next 30 years¹⁴. The downsizing and closure of state hospitals over the last few decades has resulted in trans-institutionalization of SPMI patients into nursing homes and other less restrictive environments, such as boarding care homes, assisted living and other forms of community-based living arrangements. Currently, over 89% of all institutionally-based older adults with SPMI reside in nursing homes¹⁵. It is unlikely that nursing homes will be a principal resource for care of older patients with SPMI, as further effects of nursing home reform (OBRA-87) and managed Medicaid are reinforced by patient preference¹⁴.

Mental health services for the elderly SPMI population has been provided largely through community mental health centers (CMHCs). The CMHCs have not been particularly attuned to, or capable of, accommodating the unique and complex needs of the elderly, and they may not be capable of coordinating medical-psychiatric treatments¹⁶. Older persons with severe mental illness also receive services from home health agencies that provide limited mental health care and, to a lesser extent, from the general medical sector¹⁴. As with patients with less severe mental illness, older patients with SPMI require close collaboration among providers in the general and specialty mental health sectors. Promising models of integrated care include co-location of medical and mental health providers, multidisciplinary treatment teams and cross-training of medical-psychiatric providers. These programs must also include social support services to maintain function and improve quality of life, integrative case management

services, home-based residential family support services, caregiver training and psychosocial rehabilitation¹⁴. Managed care may be the vehicle to promote such service integration because of the possibility of pooling resources from federal, state and local funding agencies. Appropriate risk adjustment mechanisms to account for the psychiatric medical and social service complexity of these patients will be required for programs to be successful.

Nursing Homes

In 1997 almost 1.5 million elderly resided in nursing homes. One-half of these people were aged 85+ and three-quarters were women¹⁷. Nursing homes have supplanted state hospitals as the major loci of institutionally-based long-term care for older adults with psychiatric disorders. Surveys of nursing home residents show uniformly high prevalence rates of dementia (46-78%)¹⁸⁻²⁰, and clinically significant depression (20-40%)²¹. Early in the trans-institutionalization movement, nursing homes became the repository of many SPMI patients. Current trends, however, find many older SPMI patients live in community settings¹⁴.

OBRA-87 legislation, also known as the Nursing Home Reform Act of 1987, was enacted in response to inappropriate and inadequate care for mental illnesses in nursing homes. The legislation restricted the inappropriate use of restraints, physical and pharmacologic, and required pre-admission screening for all persons suspected of having a serious mental illness. Screening was designed to improve treatment and psychosocial assessment for nursing home residents with mental disorders. In 1998, the Institute of Medicine (IOM) convened a follow-up analysis examining the effectiveness of the original legislation. From a psychiatric services perspective, the results have been mixed²². Pharmacoeconomic evidence has shown a downward trend in the use of psychotropic medications, and interventional trials designed to reduce physical-chemical restraints in nursing homes demonstrated that educational efforts complementing consultation by skilled mental health professionals had the best results^{23,24}. Physician prescribing practices have also changed. A new generation of psychotropic medications are now commonly being prescribed that have fewer side effects and are better tolerated by frail nursing home residents²⁵. Multidisciplinary treatment guidelines have been developed to deal with difficult psychiatric and behavioral problems, such as depression and agitation, in dementia patients. Less certain outcomes of OBRA-87 include unnecessary tensions between the legitimate use of medications and federal/state survey procedures; collection of uniform information on nursing home residents that do not have sufficient flexibility to measure quality-of-life outcomes or quality indicators; logistic barriers for medically necessary psychiatric services; and the unintended effect of establishing incentives for the inappropriate provision of some mental health services, e.g. psychotherapy services for severely demented patients²².

Important components in state-of-the-art mental health services in nursing homes include "intrinsic" and "extrinsic" mental health services²². Intrinsic services refer to the biopsychosocial elements of daily patient care activities and range from the nursing home environment to individual attitudes of professional staff, which are tied to respect, dignity and empathetic interpersonal exchanges. In addition, intrinsic services may include specialized settings and discrete units that provide behaviorally orientated services with highly trained staff. Dementia care units, "special care units" and psychiatric nursing home units are examples of specialized intrinsic services that are relevant to the treatment of older persons with mental disorders. Extrinsic services are linked to the ability of nursing home residents to gain access to specialized psychiatric services in a timely, efficient and sensitive manner. Extrinsic services generally refer to

specialized mental health services that are provided from outside the facility through a consulting or other contractual arrangements. A variety of consulting models of mental health services to nursing homes have been described and optimally include components of assessment and evaluation, with a strong emphasis on collaborating with the treating medical physician and on educating nursing staff²⁶. However, surveys of nursing home directors indicate that there is considerable unmet need for psychiatric consultation services, especially addressing non-pharmacologic management and staff training. However, incentives are lacking for adequate service provision by psychiatrists in nursing homes and there are substantial challenges to identifying the most effective interventions and services²⁷. Appropriate institutional and patient outcome measures need to be developed that can identify the most cost-effective intrinsic and extrinsic mental health services for nursing homes.

Acute Inpatient Hospitalization

Geriatric patients with late-life mental disorders requiring acute inpatient hospitalization are principally and appropriately cared for in secondary and tertiary care hospitals¹. Inpatient units within general hospitals have access to subspecialist consultation and diagnostic technology required to provide accurate diagnosis and treatment recommendations. Over the last 15 years, there has been a substantial increase in the rate of admissions of geriatric patients to nine federal general hospitals. Specialized medical psychiatry units have increased the levels of sophisticated treatment for patients with severe mood disorders, mood disorders complicated by psychotic features, and those elder patients with mixed medical and psychiatric disorders¹. Inpatient services offer multidisciplinary interventions, including psychiatric services, family evaluation and therapy, social service evaluations and, in ideal situations, coordinated aftercare services.

FINANCING MENTAL HEALTH CARE: THE UNDERPINNINGS OF THE CURRENT STRUCTURE OF THE US SYSTEM OF CARE

Ideally, health systems follow the rule that “form follows function”. In the US system of care, a more cautious approach might be “form follows finance”. In this respect, the character and dimensions of mental health services for older adults in the USA has flowed directly from the structure, incentives and limitations of the system of financing and reimbursement. In this section we will describe these recent developments in fee-for-service financing of mental health services for older adults, followed by a discussion of current trends in managed care. For geriatric patients with late-life mental disorders, Medicare is the principal payment source for acute psychiatric services in the USA. Aside from out-of-pocket expenses, state-managed Medicaid, a blended Federal and state insurance program for the poor, is the primary source of payment for institutional and long-term care services. Hence, we will concentrate on an overview of Medicare and Medicaid as the two principal sources of payment for mental healthcare services provided to older persons in the USA.

Traditional Fee-for-service Medicare

Medicare, the federally funded health insurance program, is the primary payer of acute general health and psychiatric care services for the elderly, people with chronic disabilities and people with chronic renal failure. In 1997, approximately 39 million individuals were covered by Medicare, of whom about 33.6 million were

aged 65+. Total Medicare expenditures in 1997 were almost \$207 billion and accounted for more than 11% of the US federal budget²⁸. Medicare’s nearly universal coverage for the elderly is important because of the impact of adverse risk selection on insurance premium costs. Adverse risk selection refers to the attraction of high-cost consumers to insurance plans that offer coverage for high-cost conditions. In this respect, insurance plans that cover high-risk populations (e.g. the elderly, with multiple comorbidities and chronic conditions) are likely to assume disproportionate risk compared to insurers covering services of younger populations with low use of expensive services such as acute hospitalizations and long-term care. In other words, the actuarial risk for high medical service utilization among the elderly is high. Thus, if Medicare were privatized and premiums reflected actual utilization, costs would be prohibitive for most older adults. This effect would be exaggerated for elders, as 11% of them live in poverty and another 6.4% are between poverty and 125% of the poverty level^{28,29}.

Traditional Medicare is similar to typical indemnity insurance products featuring retrospective fee-for-service (FFS) payment, deductibles and co-insurance, but it does not have limits on annual personal spending. It also does not fully cover medical equipment costs, and fails to cover prescription medicines and the costs of long-term care²⁸. Cost sharing and uncovered benefits have created the private “supplemental insurance” market, the premiums for which constitute the largest source of personal spending for community-dwelling beneficiaries²⁸. Supplemental policies may have inpatient and outpatient mental health benefits, designed to cover co-payments and deductibles. They do not alter basic coverage limits.

The proportion of Medicare expenditures devoted to mental health is relatively small, however. For example, in 1996 Medicare expended about \$9.8 billion for mental health services, up from just under \$5.1 billion in 1994^{30,31}. Most Medicare expenditures for mental health services are for Part A services, and less than one-half of 1% are for older adults in non-institutional settings^{32,33}.

Medicare’s Benefit Design for Mental Health Services: Fee-for-service and Managed Care Arrangements

Traditional FFS Medicare has two components; part A, which covers inpatient psychiatric hospital care (up to 190 days life-time maximum in free-standing psychiatric hospitals and unlimited days in general hospital psychiatric units) and outpatient care in some hospital-based clinics and other hospital technical fees³⁴; part B covers medically necessary physician, partial hospitalization and related ancillary services. Psychotherapy services provided by psychiatrists, non-psychiatric physicians, psychologists and other mental health providers, as well as outpatient electroconvulsive therapy are subject to a 50% co-payment, while medical management services are subject to a 20% co-payment³⁴.

Since the enactment of the Medicare legislation in 1965, reimbursement policies have been a financial barrier to accessing needed mental health services for the elderly and disabled, especially for outpatient services. Until reforms were enacted in the late 1980s, Medicare’s inpatient coverage was similar to private insurance, while outpatient service coverage was de minimus³⁵. For example, Medicare reimburses inpatient services carefully, less a 1 day deductible for the first 60 days; it requires a 25% co-payment for days 61–90 and a 50% co-payment for days 90–150³⁴. In contrast, in 1966–88 Medicare covered outpatient mental health services up to a maximum of \$500, subject to a 50% co-payment, e.g. Medicare only paid \$250. In 1984, limitations on medically-based psychiatric services for Alzheimer’s disease were not subject to the \$500 and 50% cap. The Omnibus Budget

Reconciliation Acts of 1987 and 1989 (OBRA-87 and OBRA-89, respectively) changed reimbursement for outpatient psychotherapy services. OBRA-87 raised the \$500 cap for psychotherapy reimbursement to \$2200, but retained the 50% co-payment, effectively paying only \$1100. OBRA-87 exempted medical management of psychotropic medications from the limit, in addition to reducing the co-payment to 20%. Partial hospitalization services were authorized. OBRA-89 removed the cap on outpatient mental health services, although the 50% co-payment was retained³⁵.

Changes in benefit design contributed to increasing expenditures for mental health services by 136% between 1987–1992³⁶. Correspondingly, service utilization increased, thus correcting the historic underutilization of mental health services by the elderly, e.g. mental health service users rose 76%, and the number of services per beneficiary over age 65 years rose 15%³⁶. Difficult to estimate, however, are expenditures for mental health services delivered by physicians in general medical settings, which reflect actual treatment services for psychiatric conditions not coded by providers or treatment services for psychiatric conditions misdiagnosed as general medical disorders.

Although increases in expenditures for mental health services provided to older persons over the last decade suggest that progress has been made in better meeting the need, it is important to note that most of the services remain biased towards costly inpatient care. In 1994 about 12.7% of Medicare claimants had a MH/SA disorder based on primary diagnosis and/or procedure codes³⁷. Cano *et al.*³⁸ estimated that in 1995 Medicare beneficiaries aged 65+ with a primary psychiatric diagnosis accounted for 325 000 hospital and skilled nursing facility stays and accounted for approximately \$1.8 billion or 53% of all acute psychiatric payments made by Medicare. The majority of admissions were to psychiatric units in general hospitals (42%), followed by general hospital admissions (29%), psychiatric hospitals (15%), and skilled nursing facilities (14%). Overall, the burden of mental disorders in the elderly is substantial; Smyer and Shea³⁹ estimated that the total direct costs for mental illness for individuals aged 65+ were \$17.3 billion.

Medicaid

Medicaid is the primary public insurer for acute care for medically indigent populations and for long-term care in the USA. Medicaid is a joint federal and state program, with an individual state contributing up to 50% of costs. Variability in benefit design among states makes it difficult to generalize about the effects of Medicaid on care nationally¹⁴. For example, states differ on eligibility criteria (this applies to people eligible for both Medicare and Medicaid), on the scope of coverage for inpatient and outpatient mental health services, pharmacy benefits, co-payment arrangements for enrollees, pre-authorization rules for inpatient and outpatient services, and managed care arrangements. Common to Medicaid programs, however, is a 20–30% reduction in reimbursement schedules compared to regional market rates¹⁴.

Figure 127.1 summarizes national Medicaid expenditures in 1995. Excluding administrative expenses and disproportionate share allocations to hospitals serving large numbers of poor people, Medicaid spent about \$132 billion in 1995 on about 34.8 million recipients, of whom only 11% were aged 65+. However, this latter group accounted for about 30% of all expenditures. About 19% of Medicaid expenditures were spent on nursing facilities, 6% on home health services, and only 2% on mental health services⁴⁰. It is difficult to determine what proportion of the \$7.1 billion in Medicaid expenditures for mental health services in 1994 was for older adults³¹. Medicaid spent approximately

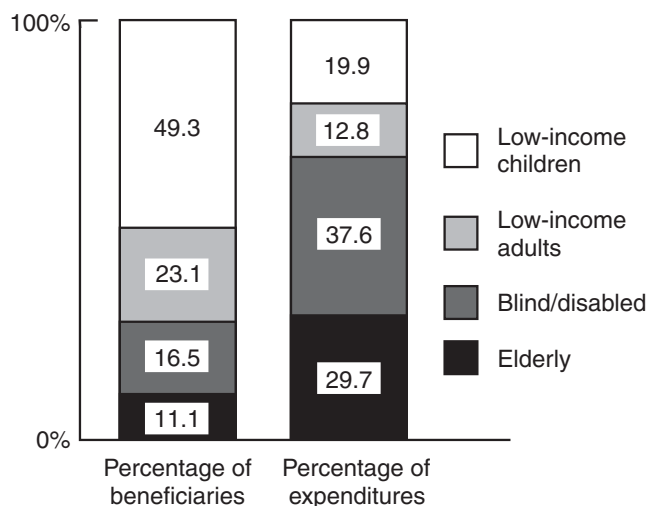


Figure 127.1 Distribution of Medicaid beneficiaries and expenditures in 1995: \$132.3 billion for 34.8 million recipients. Adapted from MedPac

\$10 129 per elderly beneficiary, and 75% of this amount was allocated to long-term care services⁴⁰.

Mental Health Managed Care for Older Adults

Over the last decade, managed care has had stunning impact on private commercial and public financing and delivery of mental services in the USA. While universal definitions of managed care have not been agreed upon, for the purposes of this chapter, managed care is defined as systems of care that integrate the financing and delivery of appropriate healthcare services to health plan enrollees by means of provider network arrangements. Managed care organizations (MCOs) furnish comprehensive healthcare services; set standards for the selection of providers; use formal and ongoing quality improvement and utilization review programs; and place emphasis on preventive services in order to avoid more costly medical care services. MCOs use incentives to use health plan providers and services in order to limit out-of-network providers⁴¹. Recall this definition of managed care later in the chapter, when we discuss demonstration projects designed to integrate geriatric healthcare services for the frail elderly.

To a large extent, managed care has succeeded in reducing healthcare expenditures in both the general health sector and specialty mental health sector, especially for private commercial health plans. For example, under the Federal Prospective Payment System (PPS) to hospitals, inflation in global operating costs per hospital case declined from a yearly average of 9.5% during 1985–1990, to a yearly average of –0.5% during 1993–1997⁴². The rate of growth of global physician expenditures has also declined. Using the Medicare Economic Index (MEI), which measures various inputs used to produce physicians' services, such as earnings, staff salaries, supplies, etc., MEI increases in 1985–1992 averaged 3.1%/year, but have declined to 2.1%/year since 1992⁴³. In the non-Medicare specialty mental health sector, cost reductions have also occurred. The Hay Group Management Corporation and the National Association of Psychiatric Hospital Systems (NAPHS) recently reported that the value of behavioral healthcare expenditures for commercial insurance plans (non-governmental insurance) decreased in 1988–1997 by 54.1%, compared to 7.4% for general healthcare costs⁴⁴. As a proportion of total healthcare benefit costs, behavioral health benefits

decreased from 6.1% in 1988 to 3.1% in 1997. The disproportionate reduction in behavioral health benefit expenditures can be attributed to both impositions on utilization patterns (24.6% decline in outpatient utilization) and benefit design (57% of plans imposed day limits on inpatient care, and 48% placed outpatient visit limits by 1997)⁴⁴.

Medicare Managed Care and the Balanced Budget Act of 1997

Encouraged by the private sector's success in managed care, coupled with rising healthcare costs, federal initiatives have stimulated the growth of Medicare managed care, especially since the enactment of Public Law 105-33, the Balanced Budget Act of 1997 (BBA-97). According to the Department of Health and Human Services' Health Care Financing Administration (HCFA), about 6.1 million Medicare beneficiaries were enrolled in Medicare risk-managed care plans by December, 1998⁴⁵. The enrollment growth has slowed in 1998–1999 and Health Maintenance Organizations (HMOs) holding nearly 100 risk contracts have indicated that they will withdraw from Medicare managed care in 1999 (about 409 000 enrollees) because of payment rates and regulatory burdens⁴⁵. Medicare predicts that about 44.5 million Medicare beneficiaries will be in managed care programs by 2008⁴⁶.

Several managed care options existed for Medicare patients prior to the BBA-97. These included Medicare risk contracting (MRC) plans, point of service options (POS), social HMOs or demonstration projects called programs of all-inclusive care for the elderly (PACE). Social HMOs or PACE programs are demonstration projects that combine Medicare and Medicaid funding into one funding base, providing a continuum of healthcare services, including inpatient, outpatient and long-term care⁵. The BBA-97 created the Medicare+Choice program, a new Part C of Medicare. Medicare+Choice expanded these options to include medical savings accounts, POS options that allow patients to select from a broader panel of practitioners outside of the HMO network, religious fraternal benefits plans, and other coordinated care plans meeting a set of established standards⁴⁶.

MRC plans are the most frequent type of Medicare managed care arrangement. A MRC plan receives a set payment per month per patient, based on a county level adjusted average per capita cost (AAPCC), as determined by HCFA. The BBA-97 made significant changes in the payment methodology to MRC plans that have had historically high AAPCC payment rates. Beginning in 2000, the risk adjustment methodology used to pay many HMOs also changed, further reducing HMO payment rates⁴⁷. To remain solvent, MRC plans must ensure that their costs do not exceed the AAPCC payments. Concerns over the payment methodology have forced some plans to leave the market, while others have reduced benefits and increased co-payments and premiums.

By law, MRCs must offer basic Medicare benefits, including mental health benefits. In order to entice Medicare beneficiaries into joining Medicare HMOs, many plans offer enhanced supplemental benefits, such as prescription drug benefits, dental coverage, optical or hearing services⁴⁷. Neither HCFA nor the managed care industry has established policies or procedures for how mental health services should be delivered to the elderly. Largely undefined, or using criteria from commercial managed care plans, are medical necessity criteria, co-payment policies, credentialing standards for providers, hospital network standards, geographic access rules, referral mechanisms and quality improvement mechanisms⁵. Case management guidelines, coordinated care for patients with medical–psychiatric co-morbidity, dementia care and long-term care policies (beyond the 90 day rule for

skilled nursing home care) have not yet been clarified. These plans may also fail to risk-adjust for chronicity and medical–psychiatric co-morbidity of late-life mental disorders, such as Alzheimer's disease or recurrent major depression³⁵.

Managed Medicaid

Over the last several years, many states have created Medicaid managed care arrangements. As of 1996, about 38.6% of all Medicaid beneficiaries were under Medicaid managed care arrangement, and increased 12% during 1995–1996⁴⁰. The move to Medicaid managed care has been encouraged by program waivers from the Federal Government under Section 1915(b) and Section 1115 of the Social Security Act⁴⁰. Section 1915(b) allowed states to mandate enrollment into managed care programs. Section 1115(a) allowed the US Secretary of Health and Human Services to approve time-limited demonstration projects that test and evaluate innovative approaches to delivery and financing of health care. Section 1115(a) also allowed some states to expand Medicaid eligibility for acute care services; however, it has been used to enroll Medicaid beneficiaries into prepaid managed care programs⁴⁰. The impact of these new financial arrangements on access to mental health and long-term care services by older persons is yet to be determined. The degree to which state-run Medicaid programs will reallocate resources away from long-term care programs used by elders towards children and low-income families is also unclear.

Mental Health Managed Care Arrangements: The "Carved-out" vs. "Carved-in" Debate

Most Medicare MRC plans "carve out" the mental health benefit package, similar to what they do for commercial patient populations. Mental health "carve-outs" refer to the practice of setting aside funds for mental health benefits and then contracting with a vendor, who is responsible for managing all mental health services. Benefits of carve-out mechanisms include protection of the HMO from expenditures above the contracted percentage of premium paid to the carve-out vendor, reduction of HMO staffing and space needs, patient confidentiality and mental health professional input for difficult cases. Proponents of carve-outs purport that these financial arrangements for mental health services are superior because vendors are able to manage costs and services through superior technical knowledge, skills and service delivery networks⁵⁸. "Carve-out" approaches to Medicare managed care may not be the best approach for older adults because of medical–psychiatric co-morbidities. These arrangements may not provide coordinated psychiatric and primary care services⁴⁸; may restrict physician involvement through profiling and paneling⁴⁹; may increase access barriers to care⁵⁰; and may limit the quantity of needed services⁵¹. Carve-out arrangements may pose problems with demonstrating cost offset for mental health services, especially for those patients with significant medical–psychiatric co-morbidity¹⁴. A study of dominant carve-outs providing service to MRC payers demonstrated little or no requirement for specialty-trained providers or evidence of experience in caring for the elderly⁵².

Some argue that mental health benefits for older adults should be "carved-in", e.g. where mental health services are included as part of the general health benefit. Advocates argue that this benefit design better integrates mental and physical care, decreases access barriers due to stigma, and produces cost offsets in general healthcare expenditures because of the high medical–psychiatric co-morbidity among older adults¹⁴. Other benefits include better collaboration among psychiatric and physical health providers, as

Table 127.1 Selected features of managed care programs for the frail elderly: PACE, social health maintenance organizations (S/HMOs), and EverCare programs

Characteristic	PACE	S/HMO	EverCare
Setting	Community setting	Community setting	Nursing home
Start-up	1971, On Lok Program 1990, PACE	1985, first generation of S/HMO 1997, second generation of S/HMO	1994
Number of active programs in 1999	25	First generation, 3 Second generation, 1	Six demonstration projects
Focus	Integrate delivery and financing of acute and long-term care for frail elderly	First generation: test models of integrating acute and long-term care, and social services in a capitated HMO setting Second generation: required improvements in service, and benefit design including: geriatric assessments, multidisciplinary teams, expanded case management services for individuals at risk for disability	Enrolls permanent nursing home residents into managed care programs with a focus on providing Medicare covered outpatient services in order to reduce hospital and emergency room use
Eligibility requirements	Eligible enrollees must meet state nursing home eligibility requirements	Initially limited to frail beneficiaries aged 65+	Nursing home residents
Benefit structure	Acute and long-term care benefits covered through Medicare, Medicaid and private capitation payments. Prescription drugs are covered	All Medicare benefits, expanded benefits (similar to typical Medicare Risk Contracts), and long-term care benefits. Prescription drugs are covered	Similar to Medicare Risk Contract but no prescription drug benefit
Enhanced mental health services	Not addressed	Not addressed	Not addressed

Adapted from MedPac⁵⁶.

well as improved psychiatric services for those older patients who receive most of their mental health service from primary care providers. Carved-in benefit designs are not without hazards. Functional integration of mental and general health services are far from guaranteed, and mental health services are likely to receive low priority in these types of managed care arrangements. Comprehensive services for mental health, e.g. parity, may not occur, and payment methodology that adequately risk-adjusts for the more seriously ill patients are not well developed, thus placing a health plan at risk for substantial financial loss^{14,53}.

EMERGING AND FUTURE TRENDS IN INTEGRATED SERVICES AND FINANCING

While recent legislative changes in Medicare and Medicaid have incentivized some efforts toward the goal of a seamless continuum of care for the most frail and vulnerable geriatric patient populations, much work remains. Promising demonstration projects have emerged, such as out-of-pocket-financed community-based dementia respite care programs⁵⁴. While Medicare's psychiatric and mental health benefit has been liberalized over the last 13 years, parity between mental health and general health care benefit coverage has not occurred. Nevertheless, the BBA-97 expanded three demonstration projects designed to care for those frail Medicare beneficiaries who need more long-term chronic and acute care. Programs include the program of all-inclusive care for the elderly (PACE)⁵⁵, social health maintenance organizations (S/HMOs) and the Evercare Demonstration Projects care⁵⁶. These programs were not specifically designed for older adults with late-life mental disorders; however, because the frail elderly are at increased risk for psychiatric illness, mental health services will be part of the package of services.

Table 127.1 summarizes the select features with PACE, S/HMOs and the Evercare programs. PACE is a community-

based program that is designed to delay or prevent the use of hospital or nursing home care by providing a comprehensive range of preventive, acute and long-term care services in community settings. The first generation of S/HMOs was designed to test the integration of financing and service delivery of a full range of acute and long-term care services through capitated HMOs. The second-generation S/HMOs have expanded programming, including such services as geriatric assessment, case management services and a multidisciplinary team approach to care. EverCare is a program that enrolls permanent nursing home residents into managed care programs and provides "in-place" primary care services to nursing home residents⁵⁶. Primary care services include both acute and preventive services, provided within the nursing home setting.

All three programs use managed care financing, case management principles, and provide a wider array of medical and social services. Enrollees in the S/HMOs have similar demographic characteristics to those beneficiaries enrolled in Medicare, but PACE and Evercare enrollees are significantly older, and a higher proportion of them are Medicaid-eligible. Unfortunately, none of these demonstration projects have specific requirements for the provision of mental health services, although the profile of beneficiaries expected to enroll in these programs would have considerable psychiatric co-morbidity. Program evaluations for the PACE Program and S/HMOs have demonstrated mixed outcomes. For both PACE and S/HMO, enrollee satisfaction was generally high⁵⁶, although enrollees in the S/HMO with physical impairments were usually less satisfied⁵⁷. Psychiatric outcomes have not been measured to date.

SUMMARY

The last half of the twentieth century included an explosion in the size of both the older-adult population and the US healthcare system. Since the introduction of Medicare and Medicaid in the

1960s, older adults have had universal access to acute health care and their demand for services has driven the development of a highly capitalized, technologically advanced but severely fragmented system. This interaction has contributed to the extension of life expectancy, a probable improvement in life quality for many, and staggering growth of expenditure. The last two decades have been consumed by efforts to contain and reduce costs, despite continued technology growth that has had marginal effect on efficiency. Throughout this period, mental health has been, at best, a poor stepchild. While enjoying advances in science and recognition of its importance, the care of older people with late-life mental disorders has evolved services that reflect modest reimbursement schemes while attempting to meet a minimal subset of population needs. The provision of mental health services for the elderly is inextricably linked to primary care and, for the frail elderly, must attend to the problems of medical-psychiatric co-morbidity. Recent reforms in the organization and financing of health care are beginning to address co-morbidity issues through integrated acute and long-term care delivery systems, such as PACE, S/HMOs and the Evercare programs. State Medicaid managed care initiatives are also creating opportunities to reallocate support to the development of home- and community-based alternative models of care, with the goal of supporting least restrictive and less costly long-term care services. Demonstration projects must examine behavioral outcomes and analyze their effects on general health, mental health, functional capacity and quality of life. Yet to be determined is the impact of these types of programs on increasing access to medically necessary mental health services.

At the dawn of the new millennium and in the face of extraordinary projected population growth of those aged 65+, opportunity must be seized from innovation. The shift of some insurance risk to providers from payers and the government may be perilous, but it is the first opportunity to align incentives to benefit the populations we serve.

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Community-based Psychiatric Ambulatory Care: the Private Practice Model in the USA

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In the USA, ambulatory or outpatient psychiatric care of individuals in later life is provided in a variety of public and private office settings. These include publicly-financed community mental health centers, hospital-sponsored outpatient clinics or services, the offices of psychiatrists (and other mental health professionals) in private practice, health maintenance organizations (HMOs, both privately and government-funded), and others. This chapter will focus on practical aspects of providing psychiatric treatment to older Americans in office-based psychiatric private practices. We will not discuss the details of treatment in these settings, but rather the “mechanics” of the process. An important focus of community-based psychiatric ambulatory care is the need to create relationships with other community services and providers of care and assistance. This provides both a framework and a means by which many of the services are provided. Some of the barriers and obstacles to care will also be reviewed.

In providing community-based ambulatory care, the psychiatrists must have a comprehensive and patient-centered focus, following the patient to provide whatever psychiatric treatments are needed in whatever setting. While there are some geriatric psychiatrists who limit their activities to specific treatment locations, such as in offices or in-hospital programs, this type of care may require the patient to be seen and treated sequentially in many places and circumstances.

INFLUENCES ON PRIVATE PRACTICE

For the purposes of this chapter, we will define private practitioners as independently employed or self-employed psychiatrists who work alone or in small groups. These practitioners provide treatment to patients who individually seek their help, and who pay for services received, primarily with Medicare health insurance benefits. Notwithstanding the large-scale reorganization occurring in the American healthcare system, private practice remains the widespread model of medical practice in the USA. In many ways it has served as a starting point for the pattern of care provided in the other settings mentioned above. Many of the techniques discussed below are applicable to other models of treatment. Older patients seek care from private practitioners or other mental health providers with varying degrees of utilization and satisfaction^{1,2}.

The private practice of medicine in the USA has evolved as a cottage industry within an historically unsystematized, free-enterprise, fee-for-service climate. Changing lifestyle preferences

and demographics among young physicians are having some impact on this pattern. From its inception in 1965, the Medicare system has had a built-in prejudice against the provision of outpatient psychiatric services. There remains a discriminatory 50% patient co-payment requirement for all psychiatric treatment services delivered outside of an acute care hospital setting (unlike all other covered medical services, for which the patient co-payment is 20%). This has constrained provision of these services to the elderly. Over the past 5–10 years, however, much greater influence has been felt from a multitude of outside forces, at times impeding geriatric psychiatry practice^{3,4}, but in some cases expanding it. These forces (to list a few) include: efforts by the federal government to rein in Medicare spending^{5,6}; the penetration of health maintenance organizations (HMOs) into Medicare; the advent of federal nursing home reform regulations; the growing presence of healthcare agency accrediting bodies, such as the Joint Commission for the Accreditation of Healthcare Organizations (JCAHO) and the National Council of Quality Assurance (NCQA); the creation by the American Board of Psychiatry and Neurology of subspecialty board examinations in geriatric psychiatry; social attitudes (e.g. public attitudes about medical care and doctors, as well as about mental illness and psychiatric care); the influences of physician attitudes⁷ and medical malpractice litigation⁸; the availability of newer psychopharmacologic agents (including their increased acceptance by the public and their increased utilization by primary care physicians); patient finances; and folklore and “common sense” of both the professional and the general population. Geography also is a factor, as there is significant regional variation among practitioners and communities in different areas of the country. Adaptation of geriatric assessment and treatment principles has been slow, for the most part, in private medical practice. Nevertheless, successful practitioners need a working knowledge of these forces in order to function. As a relatively new field in the USA, there are few established examples of psychogeriatric care that have proven generally applicable. In fact, the fee-for-service system in the USA has thus far been a relative failure in geriatrics, since it has not incorporated many of the accepted principles of geriatric care⁹.

The incursion of managed care HMOs into the Medicare system held out promise to change this state of affairs. HMOs, in theory, employ methodologies such as integrated delivery systems, screening, prevention and case management that are ideally suited to geriatrics¹⁰. Their track record has been disappointing, however, and their approach to managing mental health care has been largely ineffective for the elderly. Managed care

companies often “carve out” the management of mental health services by subcontracting it to managed behavioral healthcare companies with specialized expertise in mental health benefits management¹¹. Such companies rarely have any expertise in geriatrics, and do not appreciate its differences from general adult psychiatry¹². These companies have often ageist attitudes built into their coverage utilization guidelines, and inappropriately limit treatment or completely deny it, especially involving members with Alzheimer’s disease, which they do not view as a covered psychiatric disorder. To the extent that managed care has penetrated Medicare, this practice has made the practice of geriatric psychiatry unnecessarily burdensome for many American psychiatrists.

PRIVATE PRACTICE AS A BUSINESS

Another important factor in discussing this type of psychiatric practice is the awareness that it is a business, and that the patients are customers. As such, it behooves the psychiatrist to organize the practice and to provide services in ways that answer the needs of these customers. The psychiatrist may help the patient to define these needs, provide information about them, alter them, or aid them in various ways. The psychiatrist may need to refuse patients’ requests when professional judgment dictates this. American consumers, especially the adult children of geriatric patients, are becoming increasingly distrustful of the healthcare system and doctors. If the psychiatrist does not do a good job or does not adequately address at least some of the patient’s needs (and/or their adult child’s needs), the psychiatrist may lose that patient’s business. Thus, it is clear that the interaction between them is an exchange of service for payment. By providing a comprehensive service, as mentioned above, the psychiatrist may provide more services in more locations, which is often very satisfying and helpful to patients. At the same time, the business opportunities for income are maximized. In a community-based private practice, people are often referred to the individual doctor, rather than to a hospital, a university or a public clinic where they may be assigned a doctor. Patients may be referred because of the doctor’s quality of service, reputation or relationships with the referring party. These qualities therefore become significant aspects of the psychiatrist’s success in business as well as clinical practice.

Important factors in satisfying the patient/customer include:

1. *Cost*: reasonable fees and/or helpfulness and knowledgeability in filling out insurance claims.
2. *Accessibility*: convenient and comfortable office location and surroundings.
3. *Availability*: the availability of the psychiatrist to go to the patient if needed (e.g. to consult at a medical hospital if the patient is admitted by another physician for a physical ailment, or to see the patient at home, in a nursing home, or an assisted living facility) is very important. The convenience of the geriatric psychiatrist going to where the patient lives, rather than the patient coming to the doctor’s office, is very attractive to family caregivers responsible for transportation. The viability of this form of practice, in private practice, depends upon arrangements with facilities that ensure an adequate volume of patient visits for each trip to the facility.
4. *Scheduling*: flexibility to see patients at convenient times without excessively long delays in scheduling appointments. This is crucial, since many frail patients are brought by their adult children who work.
5. *Communications*: the ability to contact the psychiatrist quickly and easily at need (e.g. by telephone by the patient and, when appropriate, by the patient’s family). This includes the willingness of the psychiatrist to return such phone calls quickly, and the friendliness and accuracy of the psychiatrist’s secretary or answering service. It also includes the ability and willingness of the psychiatrist to speak to the patient (and family) about his/her symptoms, illnesses and treatments in a clear and patient manner.
6. *Concern*: the feeling that the therapist has a genuine interest and concern for the patient. This feeling of concern extends to the patient’s interactions with the office staff. This is an especially vital factor for the older population¹³.
7. *Confidence*: patients need to feel that the psychiatrist knows what the patient’s problem is and has an idea about what can be done. The doctor does not need to define answers, but must indicate a grasp of the situation and some ideas for an approach to it. This helps to provide a structure to what is often a strange and frightening experience. Empathy with the patient’s distress is very helpful in this, as is reassurance to the patient that his/hers is not the worst case the doctor has ever seen (a common fantasy).

There are only limited data available on income and workload for geriatric psychiatrists as a group. For all psychiatrists, 1998 median annual gross income was \$171 490 (a 3.5% increase from 1997) and annual net income was \$118 630 (a 4.33% increase from 1997). This was the second lowest income of the 20 largest specialties (above general practitioners) surveyed by Medical Economics that year. The rate of inflation in the year 1997–1998 was 1.6%. Comparable 1998 income data for all US physicians show an annual gross income of \$256 290 (down 0.7% from 1997) and a net income of \$163 940 (up 2.2%); for non-surgical specialties the gross income was \$227 300 (up 1.7%) and the net income was \$147 140 (up 2.4%)¹⁴. When these data are compared to the median annual net income for psychiatrists in the USA in 1989, which was \$103 570 (the fourth lowest of 15 office-based specialties surveyed that year; the only doctors who made less were general practitioners, family physicians and pediatricians)¹⁵, we find that the income of psychiatrists had risen 14.5% in that period. The median net income for all fields of medicine rose just under 25% during the same period, while the cumulative inflation rate added up to 35%.

Although most American psychiatrists see few or no geriatric patients, this trend is changing somewhat. There were over 5400 out of the over 36 000 members of the American Psychiatric Association who expressed an interest in geriatrics during their 1997–1998 Professional Activities (Biographical) Survey¹⁶. The membership of the American Association for Geriatric Psychiatry has grown to over 1800¹⁷ and interest among general psychiatrists is increasing. In 1991, the American Board of Psychiatry and Neurology first administered a Board Certifying subspecialty examination in geriatric psychiatry. As of September 2000, there were 2508 individuals who have passed this examination¹⁸.

In 1996, 18% of American general psychiatrists had geriatric caseloads exceeding 20% of their practices¹⁹. Overall, in this 1996 survey of 970 responders, an average of $14.0 \pm 17.7\%$ of their psychiatric patients were aged 65 +¹⁹, compared to 8.4% found in a 1987 study²⁰. When psychiatrists who provide a higher proportion of geriatric services (more than 20% of their case load—HGP) were compared to those who were low-volume providers with the elderly (less than 20% of their workload—LPGs), it was found that the HPGs spent proportionately less time in their offices (although still spending most of their time there), more time in hospitals and significantly more time in nursing homes, than LPGs¹⁹. In view of relatively low numbers of psychiatrists with a specific interest treating the elderly, when the medical and general communities know that a particular psychiatrist is a geriatric specialist, there is usually no shortage of patients needing these services.

OFFICE PLANNING AND DESIGN^{21,22}

Establishing a practice to treat older patients requires some attention be paid to the setting in which such treatment will occur and to factors that might act as barriers to treatment. Offices that can only be reached by climbing stairs, or those with varying levels into which one must step up or down, are difficult and potentially hazardous. Long corridors that must be traversed are similarly problematic. Chairs should be available that are sturdy and have armrests and firm seats, high enough for ease in sitting or rising. Adequate lighting, readable signs and patient information literature should be planned with poor vision in mind. Area carpets, spring-hinged doors and other possible hazards should also be considered.

Mobility and transportation problems are another potential obstacle to treatment. Selection of an office location in a rural community, or in an area with poor public transportation or poor handicapped access, may be factors. Treatment may be interrupted during the winter months if the cold interferes with the patient's ability to get to the office. Some communities have senior transportation services, which will take people with limited mobility to physicians if reservations are made 24–48 hours in advance. Some hospitals may transport patients to and from the hospital or to physicians' offices located in adjoined buildings. Offices may also be located in senior retirement buildings or communities.

THE BEGINNING OF THE RELATIONSHIP

Older patients seek out, are referred to or are brought to the psychiatrist's office for care. An initial "gatekeeper" function may occur by means of inquiries (usually by telephone) into the reasons for the request to be seen, the age of the prospective patient, the referring source, the status of insurance coverage or other financial information. Such inquiries may lead a particular practitioner who prefers to specialize in geriatrics to decline to accept an adolescent as a patient, or to suggest that a patient being seen by another psychiatrist first discusses the idea of transferring with the current therapist, or to refer the patient to a geriatric psychiatrist in a geographically more convenient location.

A prospective patient, once given an appointment, should be told about additional information the psychiatrist would like to have available at the time of the first visit (e.g. the names of the patient's other treating physicians, current medications being taken, information about past psychiatrists, psychiatric medications, hospitalizations, etc.). If the referring source is a physician, family member or a member of the staff of a senior-living facility, information from them as to the nature of the problem referral may be requested.

THE RANGE OF SERVICES

Among the most important services a psychogeriatric specialist can provide are diagnostic services. Too often, inadequate or erroneous evaluation leads to inadequate or erroneous treatment. A knowledge of physiology, psychology and the illnesses of late life, a comprehensive approach to history taking, assessment and testing and the ability to formulate an appropriate treatment plan form the basis of a unique contribution by geriatric specialists^{23,24}. In fact, the ability to provide such a comprehensive evaluation and treatment perspective may be a primary reason why patients and referral sources seek the assistance of a geriatric psychiatrist.

An important aspect of the coordinated treatment plan is the collection of past information. With the patient's permission,

contact is established with the patient's family, other physicians and therapists. Past records, diagnoses, psychological testing reports, doctors' treatments, psychotherapy records, laboratory and radiological reports are collected. While not revealing confidential information, these contacts also benefit the patient by making the patient's other physician and support system aware of your activities with the patient. This increases the likelihood that you will be notified of future problems that may occur and that other medical treatments will be coordinated with you by other physicians. In the absence of such relationships, physicians may call a different psychiatrist to provide treatment, due to lack of awareness of your involvement.

As has been reviewed elsewhere^{22,25,26}, including sections of this volume, older individuals can be suitable candidates for many of the therapeutic modalities provided to younger patients, including individual, group and family psychotherapies, which utilize insight-orientated, cognitive, behavioral and other techniques. Some approaches, such as reminiscence or life-review therapy²⁷, have more specific applicability to the aging person. Modification of family therapy may be necessary, e.g. to address the role of adult children in assisting in the care of a demented or otherwise impaired parent.

Unfortunately, the federal government has been increasingly scrutinizing and denying payment for psychotherapy services for Medicare beneficiaries, including many elderly patients. Under the Clinton Administration's program "Operation Restore Trust", an effort to reduce fraud and waste in the Medicare and Medicaid programs, many psychotherapy services came to be viewed as unnecessary or fraudulent, particularly those provided to demented patients or in nursing homes^{28,29}.

Psychopharmacologic treatment of the elderly often requires alteration in the selection, dosing and scheduling of medication because of changes in absorption, distribution, metabolism, receptor sensitivity and excretion^{30,31}. Once again, the geriatric psychiatrist may be sought out in recognition of this expertise by the patient and others involved in the patient's care. Pharmacoeconomic trends in the USA create additional conflicts for geriatric psychiatrists in private practice. First, many patients who have Medicare have no prescription drug benefits (although this was an important political issue in the US presidential election of 2000). Pharmaceutical costs are escalating rapidly in the US compared to many other countries. Patients may have coverage for the physician visit but not the drugs the physician prescribes³². Those beneficiaries who have opted for managed care plans may have some drug benefits. However, treatment options are often limited by restrictive formularies designed with cost containment in mind, but often without any consideration of the greater sensitivity of elderly plan members to medication side effects. Nursing homes have also adopted formularies to contain their costs, even in the absence of significant managed care penetration. Consultant pharmacists are employed by the homes to monitor physician prescribing, with respect not only to federal regulations but also to formulary requirements. Psychiatrists in private practice are thus often under pressure to prescribe less costly drugs or to run the risk of receiving fewer referrals from primary care physicians or nursing homes.

Sometimes assistance provided may be primarily educational, such as telling the patient or his/her family about the nature of the aging process or the symptoms, prognosis and treatment of an illness. Treatment may be primarily informational, directing patients and carers to appropriate senior housing, services for the blind or hearing-impaired, continuing education programs or volunteer work. At times treatment may be directive, e.g. telling a patient to get a physical examination, buy a hearing aid or give up driving, or telling a family member to seek the assistance of respite services to provide some relief in caring for a cognitively impaired person, or to advise that the parent or sibling should no longer

live alone. The community-based psychiatrist must develop expert knowledge of the available community resources, as well as relationships with the providers of them.

The initial psychiatric diagnostic evaluation of the patient also is the time of the patient's actual evaluation of the doctor. The practitioner must address the overt and covert concerns, the anxieties and fantasies about the nature of geriatric psychiatry, the reasons why the patient is there and the treatments that will be instituted. Although these anxieties are not unique to this model, the need to address them is. Unlike treatment limited to one location or modality or situation, this relationship will be multifactorial and ongoing. Furthermore, a privately operating care provider is likely to represent an entry contact point into mental health care. If the patient and the associated significant family members are not put at ease, their questions answered and concerns addressed, the contact may quickly end.

Older patients are often novices regarding mental illness and its treatment. They are often fearful of being thought "crazy" or of being "put away". Structuring the beginning of the initial interview can help relieve their anxiety. You may start with 10–15 min of specific questions, such as address, age, date of birth, concrete information on marriages, children, parents, siblings, education, employment, interests, etc. This can also give you a lot of information in a short time, helping to give a more complete picture of the patient. Simultaneously, you are assessing aspects of mental status and memory.

Treatment of the older patient includes the time when the patient is away from the office. The patient is helped when assured of the doctor's continued interest and care. This can often be done using relatively simple techniques: (a) providing specific information tells the patient that you know what is going to happen, e.g. "This medicine is going to take 2–3 weeks to build up in your system. You may experience some side effects during that period but you will not experience the benefit for 2–3 weeks. You need to be patient during that time"; (b) assuring access and inviting communication, e.g. "My telephone number is a 24 hour number. If you have any problems or need to reach me, you can call any time". Patients rarely do call outside of office hours after being told this, but they feel very reassured; (c) specific instructions for behavior and for contacts, e.g. instead of saying, "Call me if you have any problems", saying "Call me next Tuesday", assures the patient that you want to hear from him/her. It also reduces the number of calls he/she might otherwise make before next Tuesday.

FAMILY INVOLVEMENT

Families are often interested and involved in the psychiatric care of elders. Relatives and friends can be important sources of information to the doctor. Interactions may include mediation and other interventions into the family system, re-interpretation and re-framing of past and present events, support, reassurance and education. Attention to family issues is especially important in treatment of patients with dementing disorders^{33,34}.

Because of the increased interrelationships and involvements some families have in an older patient's status and treatment, it is often vital to maintain contact and a positive rapport with the family. Also, patients will often request this. Conversely, when family members feel unnecessarily excluded or denied access to information, they can influence or disrupt the treatment entirely. This is not to imply that therapeutic confidentiality is not maintained; families generally understand this. They do, however, want to know that appropriate help is being provided. Such reassurance can have a positive therapeutic effect on the patient, as a reflection of the family's confidence in the doctor. It can also

have the practical influence/effect of helping to keep the patient in treatment.

In situations where there are no immediate relatives, non-kinship, support networks become increasingly important³⁵. The therapist may at times utilize the assistance of family surrogates in gathering information and in helping the patient. Where such networks are weak or absent, assisting the patient in their creation can be of great benefit.

RELATIONSHIPS WITH OTHER PHYSICIANS

In the absence of the formalized organization of a university environment or the planned hierarchy of a hospital or corporate structure, the geriatric psychiatrist in private practice must create or seek out relationships with other practitioners. This can be done through involvement in professional societies, participation in the activities of the community's hospitals and through non-medical social contacts. Eventually, further relationships will also be created by patients who seek psychiatric services and request that contact be established with their other treating physicians. Collegial relationships thus created can provide advice and assistance, help in monitoring the status of patients between visits to the psychiatrist and provide sources of referrals for new patients.

RELATIONSHIPS WITH OTHER PSYCHIATRISTS

As with other physicians in general, private practitioners must create a network of relationships with other psychiatrists in the area, both near and far. Those at some distance, or whose special areas of interest or expertise differ, can be sources of referrals. Other psychiatrists may receive inquiries or have patients referred to them whom they are unable to treat; they may then direct them to you. Psychiatrists who practice in closer proximity may also be sources of new patient referrals, especially when their treatment interests differ from yours. Furthermore, a certain percentage of patients, especially those with chronic or recurring illness, may be "doctor-shoppers" and spontaneously, or by referral, change from one practitioner to another over a period of time. Developing good rapport with other local psychiatrists helps in providing better care to these patients by sharing understanding of their needs, pathology and past successful treatments and by helping to avoid duplication of previously attempted unsuccessful treatments.

Formal or informal groupings of private psychiatrists may gather for continuing education and study, to help with supervision, second opinions or "risk-management" of difficult cases, or to share tasks, such as psychiatric coverage for a local hospital's emergency room. When a private practitioner takes a break, to go on vacation or to a conference, it may be one or more of these local psychiatrists who is asked to be available to take care of emergencies or to provide ongoing services to patients who are hospitalized at the time. Often such favors are done reciprocally as a courtesy.

RELATIONSHIPS WITH OTHER PROFESSIONALS

Other professional care providers with whom privately practicing geriatric psychiatrists and their patients come into contact include psychologists, social workers, nurses, speech therapists, occupational therapists, hospital administrators and the operators of nursing homes and congregate-living facilities. Knowledge of these and other community resources is essential for the geriatric psychiatrist. At times the best treatment offered to a patient may

be a referral to one of them. Needless to say, each of these can provide valuable services. As they get to know the geriatric psychiatrist, they can also be valuable resources, e.g. they can be excellent sources of information about a patient's status and functioning when the patient is not in the doctor's office. They may allow the psychiatrist to provide more and better service to patients by helping to monitor, care for and carry out treatments with the patient. They can alert the doctor when problems are developing, often earlier than the patient might have, and can assist in the management of a crisis by supporting and reassuring the patient. These individuals are sources of referral to the practitioner. They will also speak to others in the community of their experiences and contacts with the practitioner. This is an important facet of how a professional reputation is made.

RELATIONS WITH HOSPITALS

Each psychiatric program within a hospital can have its own rules, regulations, standards, patterns of practice and pattern of relations with community-based practitioners. Some hospital facilities employ psychiatrists on staff; others may not. Some programs are organized more in accord with the direction given by the hospital and the hospital-based staff. Others encourage more involvement in program planning by the community staff physicians. Some facilities are sites for training programs and have psychiatric residents who provide services. There are some geriatric facilities within free-standing psychiatric hospitals and others that are geriatric units within medical hospitals; some are located in private, for-profit hospitals, or in non-profit or public or charitably-funded institutions, or in university-affiliated programs. While the rules, staffing patterns and required paper forms may vary from hospital to hospital, these variations are, for the most part, not so onerous as to be unworkable or impossible to deal with for the community-based practitioner. In some of these settings, the community-based practitioner may be able to influence the nature of the hospital's policies and treatment program by participation in psychiatric departmental meetings and activities.

The differences among hospitals requires some flexibility on the part of the doctor, but also may allow the possibility of tailoring referrals to the hospital most appropriate to the patient. For a variety of reasons, different hospital units acquire different patient populations and characteristics. Some programs are age-segregated, with specifically designated geriatric psychiatry wards. Others are age-integrated, with younger and older patients sharing and participating in the treatment program together. Some programs may be more suitable for cognitively intact, physically healthy older people suffering from affective or anxiety disorders or relationship dysfunctions.

THE POSSIBILITY OF INPATIENT HOSPITAL TREATMENT³⁶

At the time of the initial visit or at some subsequent time in the course of the treatment of an older patient, the psychiatrist may recommend inpatient hospital treatment. The process begins with an assessment of whether the hospitalization is something that would be beneficial and therapeutic but non-emergent, or is an urgently needed admission due to imminent danger to the patient or others. Immediately after this decision, the psychiatrist must decide whether the patient is capable of consenting to this plan. Depending upon the hospitals and resources available in the community, these assessments may lead to a decision to use a particular inpatient facility. For example, there may be one that can admit people on an involuntary basis, or care for people who

are potentially suicidal or aggressive. Similar choices may result from the ability of a specific hospital's psychiatric ward to care for elderly patients who have concurrent severe medical problems, or who are wanderers, or who need the hospital's specific therapeutic approach. Other factors that affect the choice of inpatient service include locations of past hospitalizations, the hospitals used by the patient's other treating physicians, proximity to the patient and the patient's family to allow for visitation and, importantly, whether a particular hospital has a room available for the patient at the time it is needed, and whether the admission can or can not be delayed until a bed becomes available. In some cases, where room is not available locally or at the time needed or where local facilities are not appropriate, referral for hospitalization may have to be made to a psychiatrist or facility elsewhere.

ADMISSION TO HOSPITALS

Various hospitals and psychiatric facilities within hospitals may have different procedures for arranging admissions. Typically, the psychiatrist communicates with a designated person or office to make the reservation for admission. Information that must be provided at this point varies but usually consists of the patient's name, age and admitting diagnosis. Some facilities may also wish information regarding the geriatric patient's ability to function in activities of daily living, mobility, signs and symptoms of the patient that warrant admission, a preliminary treatment plan, or the likelihood of the patient being a danger to self or others.

When the patient is admitted, each hospital's usual procedure begins. Administrators and nurses fill out forms. The patient is shown to a room, belongings are put away and the staff makes the patient acquainted with the facility and program of activities. At about the same time, the private psychiatrist is notified that the patient has arrived. If not already given, initial orders are requested. When the psychiatrist is not available to come to the hospital immediately, orders might be given by telephone to the ward nurse, addressing such needs as diet, monitoring of vital signs, laboratory tests, ward therapies and medications to be started, etc.

IN-HOSPITAL TREATMENT^{36,37}

In-hospital treatments for the elderly can include the full spectrum of therapeutic approaches devised for psychiatric patients in general, although these might vary depending upon the resources and philosophy of the facility and the specific instructions of the doctor. The psychiatrist may personally provide individual psychotherapy, family therapy, psychotherapeutic medication management, electroconvulsive therapy, or other treatments, as well as ongoing diagnostic evaluation. Many older patients also benefit from group, occupational, recreational and ward milieu therapies, physical therapy, speech therapy or reality-orientation/memory-stimulating techniques. Often, the community psychiatrist is not directly involved in these treatments; the hospital's staff members provide them as part of the hospital's program and report back to the doctor regarding the patient's progress. Nurses, social workers and other staff members also inform the psychiatrist about the patient's status, symptoms, behavior and reactions to treatment as observed during the day. Coordination, mutual understanding of achievable goals, cooperation and respect between the psychiatrist and the hospital administration and staff facilitate the psychiatrist's functioning and the treatment of the patient. It is vital that a good working relationship be achieved. If it is not, the doctor can be undermined in numerous ways.

Working in the hospital requires flexibility on the part of doctors and staff. The staff must accommodate to various

physicians and their styles of treatment. The doctors must adapt to the hospital and its program, including its staffing pattern, its treatment approach and its physical plant. Patients may be seen at times under less than ideal conditions, including differing circumstances, locations, times and schedules (e.g. seeing a patient in a semi-private room, planning hospital rounds to not conflict with group therapy programs, visiting patients only to find that they are in physical therapy or getting X-rays).

POST-HOSPITAL TREATMENT

Planning for hospital follow-up begins during the hospital stay. The physician can direct the social service worker regarding possible directions and options for such problems as living situation changes, needs for at-home services, assistance or care, possible adult congregate-living facility or nursing home placement³⁸. The social worker can investigate these and coordinate planning with the physician, patient and patient's family. Other post-discharge options the doctor can order include visiting nurses, physical therapy and other home health treatments or referral to a senior day center or a partial hospitalization day program^{39,40}. The patient's needs, desires, finances and therapeutic considerations (including the options for follow-up treatment with the psychiatrist) are important factors in these choices. Similarly, the available, involved members of the family may have opinions or suggestions. They may also direct the psychiatrist's attention toward additional problems or issues they feel are significant.

An important part of discharge planning is the re-engagement of the patient in outpatient treatment in the psychiatrist's office. An appointment can be given at the time of discharge or the patient may be instructed to make an appointment within a specified period of time. The psychiatrist makes certain that needed hospital records, including discharge summary, list of discharge medications, copies of laboratory and radiograph reports and medical consultation reports, are sent to the office. This enhances completeness and continuity of care.

In addition to other usual psychotherapeutic issues that can be discussed in the post-hospital treatment, it is important to include a review of the patient's reactions to the hospital, the symptoms that necessitated the admission and the patient's progress there. Also, it is important to watch for post-hospital regressions and symptom recurrences, as the patient returns to his/her usual surroundings or to a new environment. Medication compliance and monitoring is another post-discharge task that requires attention, especially if the medication is new, if it is causing some side effects or if it requires special care in its use (e.g. special diet or times of administration).

CONCLUSION

Community-based ambulatory psychiatric care is a relatively young and growing avenue for the treatment of older adults in the USA. There have been relatively few specific models of private psychogeriatric care described to date. Aspects of the treatment of younger adults in the community are being applied to the care of seniors; however, modifications are important in order to more fully address the special problems and needs of this population.

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The Psychiatrist's Role in Linking Community Services

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In addressing the mental health needs of older adults in the USA, psychiatrists are faced with a number of significant challenges. One of them remains sheer demand: the number of patients needing services still outstrips the availability of psychiatrists, especially those with specialized geriatric training. A second challenge is that of collaborating effectively with the primary care sector who, as will be described in some detail later, provide mental health care for the vast majority of the elderly. Third is the challenge of understanding and working effectively with a wide range of non-physician providers of services to older adults. These and other difficulties will likely become more pressing as the demographic trends towards an older society are accompanied by increasing financial restrictions.

Many psychiatrists are responding to these challenges by enlarging their role from that of generalist clinicians to geriatric-specific educators, academicians and researchers, as well as sources of expertise for innovative community-based programs. We will begin by documenting the need for services, alluded to above, and follow with examples of how those role changes are being manifested in important new programs and initiatives. We conclude with comments on future directions for the psychiatrist's role in the process of change, and a discussion of some of the premises underlying the current system of mental health care for the elderly.

THE NEED FOR MENTAL HEALTH SERVICES AMONG THE COMMUNITY-BASED ELDERLY

One way to estimate the current need for mental health services among the community-dwelling elderly is to take the prevalence rates of mental illness and subtract the portion of those already receiving services. In terms of prevalence rates, the elderly appear to have rates of mental disorders about the same as those of younger adults¹. One of the more conservative estimated prevalence rates for lifetime psychiatric disorders, 12.3%², comes from the Epidemiological Catchment Area (ECA) study, while another recent community-based study documents a higher rate of 31%¹.

Using specific and fairly restrictive criteria, Shapiro *et al.*³ estimated from ECA data that some 7.8% of those aged 65+ need mental health services. If the ECA study results can be generalized to the population as a whole, and given that the vast majority (99.6%) of the nation's 29.8 million adults aged 65+ reside in the community rather than in institutional settings (US

Census Bureau statistics, 1987), then some 2.3 million community-residing elderly may be said to need mental health care.

Many of those needing services are not seeing a mental health professional. In Goldstrom *et al.*'s Bunker Hill study¹, among patients aged 65+ with a psychiatric disorder, only 42% had at least one visit to a mental health professional, compared to 68% among the 18-44 age group ($p < 0.01$) and 53% among the 44-64 age group ($p < 0.01$). Shapiro *et al.* calculated that 5.7% of the total population of elderly need, but are not receiving, mental health services. About one-third of those needing mental health services are being seen by the specialty mental health sector, another one-third receive care only from general medical care providers, and the other one-third receive no care². Recent evidence suggests that considerable barriers still exist in the public sector that prevent the elderly from receiving specialized psychiatric care⁴. The elderly have tended to be slow to report psychiatric symptoms compared to younger adults⁵.

The fact that so many patients with *mental* health needs are being seen by the *general* medical sector (the "*de facto* mental health system")⁶ is important for several reasons. Numerous studies document that primary care practitioners overlook psychiatric disorders in their patients^{7,8}, and, even when treating patients for mental disorders, tend to do so inadequately⁹. It has also been suggested that physicians may misinterpret somatic markers of depression as being due to physical illness¹⁰. In the USA, an increasing number of older adults are enrolled in managed care plans, which usually require that the primary care physician treat most uncomplicated illnesses, limiting access to specialist care. Recent evidence suggests significant differences in the treatment of older people with depression enrolled in health maintenance organizations (HMOs) compared with younger depressed patients. Older patients received fewer mental health specialty visits, fewer prescriptions for SSRI antidepressants, and were more likely to be prescribed benzodiazepines¹¹. As the managed care model expands nationwide, the evidence remains that the depressed elderly are underserved¹². Thus, there is a great need to train primary providers in the detection and treatment of mental disorders. Also, geriatric patients with mental disorders make twice as many office visits to their primary care providers as do those without a mental disorder¹. If those patients received adequate treatment, then according to the so-called "cost-offset" hypothesis, the improvement in mental health would result in substantial decreased utilization of other health care services, thus "offsetting" the expense of providing mental health care^{13,14}. Investigators estimate that treatment for mental disorders is accompanied by an overall 20% decrease in the use of general

healthcare services, especially for inpatient care^{15,16}. In a climate of fiscal restrictions, such information lends needed support to the value of adequate mental health care.

CLINICIANS AS GERIATRIC SPECIALISTS

The tremendous need for expertise in caring for geriatric patients has been part of the impetus for the evolution of geriatric psychiatry into fully-fledged subspecialty status. The American Association of Geriatric Psychiatry was founded in 1978. Recognition as a subspecialty came with the administration of the first examination for added qualifications in Geriatric Psychiatry in April 1991 by the American Board of Psychiatry and Neurology. By 1999, 2360 psychiatrists had passed the examination and there were 39 ACGME-accredited geriatric psychiatry fellowship programs nationwide.

Becoming a geriatric specialist in psychiatry involves assuming a unique constellation of familiar roles, rather than some distinctive singular role. Diversity is the hallmark of the team membership roles geriatric psychiatrists are called upon to assume. Examples of that diversity include coordinating clinical care, taking part in community initiatives and participating at various levels in educational activities.

CLINICAL ROLES

Clinical care has evolved to a model that is much more comprehensive than before, with an emphasis on multidisciplinary assessment. One successful model for a geriatric assessment clinic originating in Seattle¹⁷ included a psychiatrist, an internist and a social worker. The assessment took three to four clinic visits, one for each of the following: (a) a psychiatric evaluation; (b) a medical evaluation; (c) a home visit; and (d) a family conference at the conclusion of the evaluation, for discussion of results and recommendations. When the patient suffered from cognitive impairment, the model often included neuropsychiatric assessment.

The Seattle clinic demonstrates the emergence of a common theme that has become the standard for comprehensive geriatric assessment: a focus on the patient's psychosocial context. Families of geriatric patients have attracted much interest since the emergence of the subspecialty of geriatric psychiatry, because of the critical role they play in the care of demented elderly patients. While attention to families has always been characteristic of certain areas within psychiatry, notably child psychiatry, until the notion of comprehensive geriatric assessment was developed, families of psychiatrically ill geriatric patients were considered ancillary, rather than integral to patient care. A major catalyst behind this broadened perspective has been a large body of research, which demonstrates the vital role of psychosocial issues for the health and well-being of older adults^{18,19}. In recent years, research has focused on the mental health of caregivers and caregiver factors influencing service utilization and institutionalization^{20,21}.

Other innovative programs have been described, e.g. Bienenfeld reported on a liaison service to nursing homes²² and Reifler *et al.* described an outreach program for mentally impaired older adults in Seattle²³. However, although it is by now well established that there is a high prevalence of psychiatric morbidity in nursing homes, there remains a great need for psychiatric consultation in this setting. Legislation limits the prescription of psychotropics in US nursing homes in order to prevent the use of "chemical restraint", and pharmacists conduct periodic reviews to ensure compliance with these rules. One innovation that may enhance psychiatric involvement in the care of nursing home patients is video teleconsultation, which enables the psychiatrist to interview

patients at remote locations, usually with the nurse in attendance on the patient. Research to date suggests that this can be an effective and economically viable medium²⁴.

COMMUNITY ROLES

Social service agencies, nursing homes, local Alzheimer's Associations, home healthcare organizations, hospice care and other organizations that coordinate services for the elderly are eager for psychiatric expertise. Collaboration also allows geriatric psychiatrists to become well acquainted with available resources and to learn when to refer to such organizations. An incentive for the development of innovative community-based programs is the increasing cost of long-term care.

One example of innovative community-based care that has begun to involve geriatric psychiatrists is the adult day center (ADC) movement. The idea of ADCs actually evolved from the mental healthcare system, where day programs had been utilized for some of the more seriously ill psychiatric patients who needed greater support than could be provided by periodic outpatient clinic visits. Adaptation of that notion to the needs of geriatric patients began to appear in the mid-1970s and early 1980s, and the movement mushroomed from only 15 documented ADCs in 1975 to over 4000 by 1998²⁵.

The Robert Wood Johnson foundation initiated a national demonstration project (the Dementia Care and Respite Services Program) in the late 1980s to promote further growth of dementia-specific ADCs. One of the primary goals of the project was to determine whether centers could become financially viable through charging for their services. Centers had struggled with unstable financial bases, which depended on their ability to obtain grant support or contributions or to utilize transient state and federal funds. The 4 year program began funding for 19 model ADCs in 1988²⁶. The "Partners in Caregiving" initiative, which included the initial 19 sites and a subsequent demonstration program involving 50 sites across the USA, showed that adult day care centers could care for individuals with all degrees of dementia, from mild to severe, while remaining financially viable by meeting over 80% of their expenses through out-of-pocket payments and Medicaid²⁵.

Other community-based models involving psychiatrists have developed. Robinson²⁷ reviewed four of them, including: (a) the Channeling Demonstration, a federally-funded initiative awarded to 12 states, designed to serve severely impaired elderly people at risk of being institutionalized and to test two types of case management; (b) the Social/Health Maintenance Organization (S/HMO), a concept developed by Brandeis University, which is a managed care system of health and long-term care; (c) the On Lok Senior Services Program, a consolidated group of medical and support services based in San Francisco; and (d) Life Care Communities, which include some 600 continuing care communities nationwide, as well as the case-management delivery system called "Life Care at Home". The Channeling Demonstration, while it did not save money, did improve quality of life for clients and caregivers. The S/HMO model achieved its goal of integrating the funding and social services of long-term care, but did not succeed in integrating medical and social services or utilizing geriatric specialists²⁸. An updated series of HMOs ("S/HMO II") has begun, which have pledged to incorporate professionals with expertise in geriatrics into the range of services provided. The On Lok model, now known as the Program of All-inclusive Care for the Elderly (PACE), has been replicated in dozens of cities around the USA, with almost 3000 enrollees nationwide. Participants average 80 years of age, and have seven or eight medical conditions. All of their health care is provided by a PACE interdisciplinary team. Outcome studies are under way. So far, the

program has been shown to lower hospital and nursing home utilization costs, and most of the patients fared as well as or better than predicted, based on their baseline clinical status²⁹.

EDUCATIONAL ROLES

Many psychiatrists have responded to the acute need for geriatric-specific services and education in the community by conducting workshops and seminars for professional groups and providing consultation for various organizations. However, a response that provides for community needs for a long-term perspective is to teach geriatrics to medical students. A 1978 Institute of Medicine report urged medical schools to incorporate teaching related to aging in both the basic and clinical sciences. In 1993, the Institute of Medicine Committee on Strengthening the Geriatric Content of Medical Training convened to assess the status of geriatric training and to develop recommendations on strategies for improving the training of physicians in geriatrics³⁰. It found a decline in the number of applicants for funded fellowship programs in geriatric-related specialties. It also found that the number of geriatric faculties in all specialties nationwide was insufficient (and in psychiatry by 1221 positions) to meet the training needs of all undergraduate medical students and residents. In response to this report, the John A. Hartford Foundation initiated a project to "Integrate Geriatrics into the Subspecialties of Internal Medicine". Recognizing that the majority of the teaching of medical students and residents is performed by those in the medical subspecialties, this initiative was designed with the goal of raising awareness and competence of the medical subspecialist in the care of the elderly.

This 6 year, \$3.5 million project has sought to identify leading subspecialists, to redirect their attention to the subtleties of geriatric aspects of their discipline, and to ascertain opportunities for teaching and research within those aspects. The primary vehicle for this is the Geriatric Education Retreat (GER), a 5 day gathering of leaders in the subspecialty and geriatricians³¹. As a result of this project, nearly every national medical organization has added a symposium to their annual meeting, focusing on geriatric aspects of the discipline; fellowship training curriculum is being developed within half of the identified subspecialties; the residency Review Committee of Internal Medicine has added the requirement that all fellows must have substantial experience with patients aged 70+, and the American Board of Internal Medicine has recently adopted a "cross-content" blueprint for all examinations, which includes internal medicine. An essential curriculum in geriatric psychiatry for general internal medicine residents and geriatric medicine fellows has been described³². The Foundation has provided an extension grant to convene a GER for the specialties of Neurology and Psychiatry with the goals of "gerontologizing" the practicing neurologist and psychiatrist and infiltrating the curriculum of the residency and fellowship programs with the knowledge needed to provide competent geriatric care.

CONCLUSIONS

Several important areas of need, and examples of efforts to respond to them, have been discussed briefly in this chapter. In summary, although progress has been made in training general psychiatrists and other specialists in the care of the elderly, there remains a need for psychiatrists to improve their ability to diagnose and treat psychiatric disorders in the elderly; to participate in improved training in geropsychiatry for primary care physicians as well as for other mental health professionals; to participate in developing innovative new models of service

delivery; and finally to be willing to enter research and academic careers with a focus on geriatrics³³.

Goldman and Frank³⁴ noted that addressing the needs listed above are particularly difficult, given the emphasis on cost-containment and efficiency prevalent today, in contrast to the mood of equity and access to care that was characteristic of the 1960s. Indeed, critics of the current approach to geriatric psychiatry state that the system is problem-driven, evolving in a way that is reactive to economic, political and social forces. They argue that a prevention-driven system is needed, which springs from more proactive efforts, anticipating needs and attempting to respond to them in light of available resources.

Several premises that underlie mental health care for the elderly in the USA were set forth by Burns and Taube² and may help to explain the rationale behind the evolution of our current system: (a) there is a preference for providing care in the community instead of institutions; (b) there is a belief in the value of managed care approaches, such as the use of case managers and primary care physicians as gatekeepers to the mental health system, and organizations such as HMOs and preferred provider organizations; (c) there is a preference for service provision general providers instead of specialists when feasible; and (d) there is a belief that care should be comprehensive, continuous and of high quality.

There are, of course, many other needs that are beyond the scope of this chapter, such as those dealing with economics, health policy and political considerations. Nevertheless, there is a great potential for positive change if the basic problem areas listed above can be satisfactorily addressed.

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The Medical Psychiatry Inpatient Unit

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Medical psychiatry inpatient units primarily serving older adults have increased in popularity and numbers in North America in recent years. Several model programs have been described with respect to structure and organization, clinical care and logistical advantages in the management of older psychiatric patients who suffer from significant medical and surgical problems. The literature also suggests that medical psychiatry inpatient units offer great advantages over traditional settings in the diagnosis and treatment of elderly patients with combined disorders. Older psychiatric patients with acute medical illness, chronic medical conditions, the negative physiologic and psychologic concomitants of ageing, the problems of polypharmacy, drug interaction and compliance, together with the need for a more comprehensive and effective clinical approach, have culminated in a number of refinements in these units. The senior author's own experience in developing, organizing, operating and continually evaluating a dynamic geriatric medical psychiatry program lends support to the importance and utility of this treatment modality. This chapter will review in detail the structure, organization and clinical characteristics and financing of a medical psychiatry inpatient unit.

STRUCTURE AND ORGANIZATION OF THE GERIATRIC MEDICAL PSYCHIATRY INPATIENT UNIT

The medical psychiatry inpatient unit is generally orientated towards the admission and treatment of patients with combined medical and psychiatric illnesses. An attempt is made to integrate medical and psychiatric care, utilizing a biopsychosocial or systems treatment approach. The unit itself may be influenced by the administrative structure of the facility, the orientation of the medical and psychiatric community, the priorities among the clinical and administrative leaders and/or the general resources and expectations of the population to be served. Young and Harsch¹ alluded to three guiding principles that must be met in order for a medical psychiatry unit to succeed. These are the following: (a) the provision of a distinct type of care; (b) an improved quality of care; and (c) more efficient care. Another primary consideration in North America is the need to demonstrate that added costs to third-party payers and hospitals will yield greater benefits for both the patient population and the medical facility. The financing of these units with respect to changes in the USA are to be addressed in this chapter.

The medical psychiatry inpatient unit maintains a distinctive patient population by virtue of admission criteria. Patient

characteristics may also be determined by affiliation with various governmental or community agencies, or perhaps by other referring psychiatric facilities that are unable to provide care for medically ill psychogeriatric patients. Of course, healthier psychiatric patients may also benefit from the medical model adopted in a medical psychiatry unit. However, these units truly provide a therapeutic edge and hold promise for the successful clinical approach to a growing number of seriously medically ill or functionally compromised elderly psychiatric patients². Furthermore, the work-up and treatment may potentially be performed without major increases in length of stay. Incidentally, these units are known to provide an excellent milieu for clinical training in psychiatry and other disciplines.

The ideal physical environment of a medical psychiatry inpatient unit ensures safety for delirious or behaviourally disturbed elderly patients, facilitates the rendering of medical services, and provides a pleasant environment suitable for relatively long hospital stays. The overall space requirements do not differ markedly, but do exceed the space ideal for general hospital or free-standing psychiatric units. Presumably, the facility contains essential equipment and structure that would not otherwise be found on a psychiatric unit; the level of care delivered could, therefore, not be provided, or provided as well, on the "typical" psychiatric unit. This inpatient approach is in contrast to the consultation-liaison model of managing medically ill psychiatry patients on existing medical/surgical wards.

Features of patient rooms in a medical psychiatry unit may vary depending on the individual characteristics of the patients. Essential safety features include shatterproof windows, break-away curtain rods, electrical outlets that disconnect in response to tampering and lockable water taps, especially with semi-private baths. Certainly, for medically acute patients, lighting for bedside examination, outlets for oxygen and suction and adjustable hospital beds are necessary. A voice-call light system, a facility for glucose monitoring and availability of cardiac telemetry are highly desirable. Medical psychiatry inpatient units are also expected to provide care for patients requiring intravenous fluids/therapy or nasogastric suction, who need total care in a bedridden or debilitated state, as well as oxygen support or clinical management of common medical conditions, e.g. diabetes, hypertension, angina, congestive heart failure, chronic pulmonary disease, electrolyte or fluid balance disturbances, and urinary tract or pulmonary infections. Patients with great acuity may be too labor-intensive, i.e. the ideal unit should be designed to strike a balance between the provision of basic medical care and the provision of intensive psychotherapeutic care. The ideal unit will also consist of patient rooms that are largely private, but some

therapeutic benefits may be derived from semi-private rooms for patients who are withdrawn, or for cases in which a higher-functioning roommate can provide aid to another with respect to orientation and structure. All patient room entrances and bathroom facilities should be wheelchair-accessible. An emergency cord should be available in the bathroom and should be reachable from the floor by patients who may have fallen. The same architectural suicide prevention considerations found on general psychiatric units should be built into the medical psychiatry inpatient unit. Other needs include shower and patient-lifting devices, a bedside weight scale, suitable chairs, supply facilities/cart and, more controversially, mechanical (soft) restraints and effective methods of observation.

The unit ideally will include built-in handrails along the walls, television facilities for bedridden patients, and industrial-grade carpet to soften falls in the hallways and/or common or public areas but with vinyl to facilitate cleaning in patient rooms. Patient rooms are designed to maximize observation and minimize noise that may distract the staff. Finally, consideration must be given to whether a medical psychiatry unit will be locked, closed or controlled, and whether involuntary patients will be admitted. With adequate numbers of trained staff, it should not be necessary to have barriers to free movement. Where sufficient trained staff cannot be provided, some units have double-handed doors. The controlled unit, utilizing electronic beep or entry computer codes, ensures that wandering by cognitively impaired patients can be thoughtfully controlled. The locked or controlled-access unit provides both safety and containment. Separate areas are designed for activities and occupational or physical therapy.

The flow of staff and consultants tends to be great on a medical psychiatry unit; thus, workspace is an important consideration. Nursing units may be better divided into modular work stations with built-in cabinetry and shelving. The record and medication room can be separated by a locked door and the entire nursing station designed as an enclosed area with an observation/reception window made of safety glass. Sufficient storage space for medical equipment and supplies should also be considered. The care of complex, medical-psychiatric patients often requires frequent, small conferences among professionals of different disciplines and specialties in order to coordinate care. On-site professional offices for the medical director, social worker, head nurse and trainees afford privacy and a quiet environment for conferences and interviews. Preferably, physical therapy should be provided on-site if at all possible; a physical therapist permanently assigned to a medical psychiatric unit may incorporate psychiatric skills into the therapy treatment and may become an integral part of the treatment team. In addition to common areas for group dining, activities, educational programs, family meetings, group therapy and occupational therapy, additional space is necessary for staff meetings and conferences. An adequate staff lounge not only improves morale but also facilitates communication between the healthcare professionals and multidisciplinary personnel.

OPERATIONAL FEATURES OF THE GERIATRIC MEDICAL PSYCHIATRY INPATIENT UNIT

A multidisciplinary treatment team is generally assembled in order to address the complex needs of geriatric patients with combined disease. Each member contributes to the administration and operation of the unit, and each discipline contributes uniquely. A nurse clinical specialist with both medical and psychiatric experience may act as a milieu coordinator/supervisor for the nursing staff. Complex social work interventions require great expertise in family consultation, assessing hospital and community services and providing assistance in disposition. Consulting

psychologists with experience in neuropsychology, personality assessment and behaviour therapy are invaluable team members. The nursing staff should consist primarily of registered nurses combining medical-surgical and psychiatric skills. Practical nurses and nurses' aides can participate in the delivery of care and may also become adept in the clinical care of combined medical-psychiatric problems. Senior nurses with extensive psychiatric experience often contribute greatly to the management of behaviourally difficult patients. Irrespective of experience and background, nurses who enthusiastically support the model of combined medical-psychiatric care are ideal. Above all, the nursing staff must possess flexibility and resourcefulness combined with a practical and optimistic approach, in view of the constant changes in demands.

The usual format for the medical psychiatry inpatient unit in North America employs a model in which a psychiatrist or internist acts as an attending professional, experienced in directing a team and communicating with both mental health professionals and medical-surgical professionals. The medical director is responsible for gatekeeping, quality assurance, staff supervision, training, in-service education, trouble-shooting and the provision of consultation (directly or indirectly) in difficult cases. Liaison with community agencies, public affairs and assistance with legal and ethical problems in patient care are also required of the medical director. Administrative interfaces with the affiliated department, hospital, community mental health center and referring resources are essential. These interactions will vary according to the individual orientation and characteristics of the facility. Similarly, other third-party, legal and governmental interactions will vary accordingly.

CLINICAL AND PATIENT CHARACTERISTICS OF A GERIATRIC MEDICAL PSYCHIATRY UNIT

A variety of patient subtypes may be considered appropriate for an inpatient medical psychiatry unit. Three patient subtypes have been identified by Stoudemire and Fogel³: (a) patients with severe medical illness requiring daily medical coverage in addition to treatment of psychiatric illness; (b) psychiatrically disordered patients who require frequent but not daily medical attention from a surgeon or internist, e.g. diabetic or post-surgical cases; and (c) patients who merely require an initial consultation or periodic access or review/adjustment of medications by a medical consultant. Depression in the medically ill, chronic schizophrenia with concurrent physical disease, i.e. stroke, epilepsy, Parkinson's disease, head trauma, delirium, complications of dementia, or paroxysmal behaviour disorders, and a variety of other general medical illnesses with complex family or psychosocial issues, characterize appropriate cases for admission and treatment.

The average length of stay on a typical unit is generally approximately 7-21 days. Site-specific considerations pertaining to the selection of appropriate patients must be analyzed in order to determine specific inclusion and exclusion criteria. A well-defined priority system can then be established in order to determine which patients are admitted or provided with access to care. Regarding admission, patients on affiliated medical/surgical wards, and cases seen by the emergency or staff physicians, are typically given special consideration. Above all, patient acuity, dispositional resources and the staffing will determine the answers to many gate-keeping decisions. Moreover, the primary mission of these units remains the provision of psychiatric care to individuals who also have significant medical/surgical illness. Again, these patients are often *not* welcome in either traditional psychiatry or medical/surgical units. Finally, decisions regarding admission or transfer of inpatients from medical/surgical units may depend on the likelihood that the patient will receive true benefit, respond to

psychiatric treatment or require close observation in order to achieve diagnostic or therapeutic results.

Patients with general medical problems associated with psychiatric disturbances are challenging with respect to clinical management. However, several studies have implied that patients treated on such a unit benefit with respect to improved quality of care and shorter length of stay⁴. Medical psychiatry units are also more likely, compared to traditional units, to benefit patients with functional or cognitive impairment. For example, patients with coexisting dementia and depression are much more effectively addressed in a shorter time frame, with resultant decrease in length of stays, compared to those patients treated on a general medical unit^{5,6}. Moreover, treatment in the medical psychiatry unit with aggressive outpatient follow-up may obviate the need for nursing home placement⁶. Primary interventions on a medical psychiatry unit may include pharmacologic or somatic therapies, as well as ongoing efforts to optimize the patients' general medical/physiologic status. Coping with illness, issues relating to loss and self-esteem, feelings toward a caregiver and social changes are other important psychotherapeutic themes commonly addressed in a geriatric medical psychiatry unit. Medical emergencies, e.g. cardiac arrest or status epilepticus, must be considered and anticipated. Emergent symptoms of delirium may also need attention and in these cases, the legal doctrine of implied consent applies.

PROBLEMS, ADVANTAGES AND CAVEATS WITH GERIATRIC MEDICAL PSYCHIATRY INPATIENT UNITS

The quintessential factor that determines the success of a medical psychiatry inpatient unit is *nursing care*. Ideally, nurses recruited from medical/surgical units or critical care areas who are keenly interested in psychological dimensions of patient care, are considered best-suited to the medical psychiatry inpatient unit. Professional attire and more traditional uniforms often help to minimize confusion about the medical role of the unit. Non-psychiatric nurses may be useful in order to update and maintain medical skills of the staff, and serve to maintain familiarity with newer techniques of intravenous therapy, oxygen therapy, suction and skills in cardiac monitoring, respiratory care and postsurgical care. Because nursing is the critical element in the successful operation of these units, appropriate head nurses are invaluable and a difficult resource to obtain.

Reimbursement and financial integrity are long-standing issues with the medical psychiatry unit⁷. For example, in the USA, the Tax Equity and Fiscal Responsibility Act (TEFRA) of 1982 has determined payment for services in psychiatric units. The Balanced Budget Act (BBA) was passed by Congress in 1997 and has replaced the TEFRA of 1982. This has resulted in a reduction of approximately 7–8% in revenues for care provided in a psychiatric unit. However, patients with concurrent medical and psychiatric illness may add an average of 40% to the annual cost of health care when compared to patients without concurrent medical and psychiatric illness^{8–10}. Thus, psychiatric programs in general hospitals catering to the medically ill psychiatrically disordered patient are progressively less able to meet financial goals, because of the costs of ancillary medical service utilization, and free-standing psychiatric hospitals are also unable to include these patients in their patient mix. On average, psychiatric facilities are already losing money when they treat patients with combined medical-psychiatric illness¹¹. Further, under the Balanced Budget Act, facilities in the USA stand to lose an additional 12% or more on such cases.¹²

Medical psychiatry units in the USA that pride themselves on the ability to treat complicated geriatric patients with co-

morbid illness must consider the need to take immediate steps that will allow them to remain fiscally solvent. Further limits on admissions with no medical co-morbidity, or preference to patients with low psychiatric acuity, are the easiest solutions. However, these solutions do not allow the medical psychiatry inpatient unit to respond to the clinical needs of the complicated, high-cost co-morbid patient. A more satisfactory solution to this problem has been suggested by Goldberg and Kathol¹¹. Specifically, a partnership among medical/surgical professionals, hospital administration and psychiatric departments is suggested. A model of providing full psychiatric care, yet billing to general medical reimbursers, is recommended, allowing higher reimbursement of *per diems* through medical service billings that are adequate to cover the costs of medical tests and procedures typically not included in psychiatric or behavioural health payments. Such service integration can be done in such a way that units can actually cover direct costs and make significant contributions to the indirect costs in healthcare systems, while improving care¹³. This is particularly true if general medical patients at high risk for, or demonstrating, psychiatric co-morbidity with high healthcare utilization are targeted for admission. When this is done, costs savings for such patients accomplished through shortened length of stay can be as much as \$4000 per admission^{11,13}. Thus, it is possible to capitalize on the relatively higher reimbursement available with general medical admissions—even under the DRG system, which is also affected by the BBA, while continuing to address psychiatric difficulties.

The interaction of medical and psychiatric illness requires a unit organized in the fashion previously outlined. As forementioned, this approach allows treatment that assists the patient in moving toward recovery, with the development of policies and standards that document quality, improved outcomes and better attention for patients treated with medical psychiatry morbidity¹⁴. Under this format, the psychiatric unit director or consultant becomes involved in creating and finding ways to enforce these standards and adequate reimbursement to cover costs is achievable. Clearly, general hospital units will not be able to afford to provide psychiatric care for the medically ill, and medical psychiatric units under pressure from the Balanced Budget Act in the USA will require reorganization in order to contend with the facets of the prospective payment system introduced in 1999. Without a medical psychiatry approach, medical psychiatry patients will be scattered about the hospital, with practices that will necessarily lead to poorer quality and higher costs in the average medical setting. Hospital administrations in the USA will find that the impact on medical length of stay will create worse financial liabilities for the DRG reimbursement. Administrators therefore must continuously monitor length of stay, admission diagnoses, and dispositional plans, while maintaining a favorable prospective payment format. Direct care costs, indirect operating costs, recovery of costs, costs offset with reduced utilization, as well as other indicators of cost effectiveness, should be considered in the overall economic equation.² Mumford *et al.*¹⁵ have reported that treatment in a medical psychiatry inpatient unit may resolve problems that might otherwise become chronic and more expensive. Generally, however, units must deny admission to certain patient types whose care would exceed the permissible stay, and units must not become a way-station for problem patients. Without strict and clear guidelines, the medical psychiatry unit may become clogged and ultimately result in denial of care to a more appropriate patient. Cost-effectiveness studies are clearly needed to clarify many of these fiscal issues.

Kathol¹³ is of the opinion that many advantages of a medical psychiatry unit depend heavily upon the medical director. Perhaps the most convenient model employs collaboration between the medical director and a liaison internist or psychiatrist who

provides appropriate consultative care. Because the medical psychiatry unit as a treatment modality may increase the risk for loss of continuity of care or actual loss of follow-up, strategies for interim and longitudinal care should be carefully considered. A mechanism should be devised by the medical director for coordination of all clinical care following discharge. A distinct advantage of continuity of care after discharge from these units is the opportunity to assess the impact of psychological and psychosocial interventions among general medical populations.

CONCLUSION

The geriatric medical psychiatry inpatient unit could continue to be increasingly important as disproportionate growth occurs among the older population in North America (and Europe). An integrative model is essential in the delivery of care, preferably in a well-equipped setting that is organized to concentrate on the combined care of medical-psychiatric patients. These units are also quite useful in teaching and research programs, or in settings where large consultation-liaison psychiatry programs exist. Leadership provided by the medical director and head nurse, clearly articulated admission criteria and a well-organized multidisciplinary approach are all essential ingredients for a successful operation. Finally, as economic, social and scientific factors converge and shape geriatric psychiatry, the medical inpatient unit may prove to be an optimal setting for intensive, inpatient geriatric treatment.

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The Psychiatrist in the Nursing Home

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In the USA, nursing homes and other related long-term care facilities care for approximately 1.5 million persons annually. There are nearly 600 000 beds in these facilities across the country. While at any given time only 5% of the nation's elderly reside in these settings, up to 50% of citizens can expect to spend some portion of their lives there¹. Over the past several years, many authors have described the impressive prevalence and vast array of psychiatric disorders complicating the care of nursing home residents. When dementia is included, rates of diagnosable mental illnesses have exceeded 80%²⁻⁹. The widespread use of psychoactive medications^{10,11} and mechanical restraints^{12,13} for the treatment of disturbed behavior in this setting has been well documented⁵. The National Medical Expenditures Survey in 1987 reported that 31% of nursing home residents had a non-dementia-related primary or secondary diagnosis of mental illness¹⁴. The principal psychiatric conditions that are especially noteworthy in the nursing home include dementia-associated behavioral complications, such as agitation¹⁵, depression¹⁶⁻¹⁸, anxiety¹⁹⁻²¹, sleep impairment^{22,23,29-31}, psychosis^{19,24-26} and substance abuse²⁷. While the prominence of mental disorders in long-term care settings is now beyond dispute, in the USA, staffing patterns, staff expertise, environmental design and models of care delivery much more closely approximate subacute and chronic general medical care capabilities than thoughtfully conceived mental health services⁹. As a result, there is a great need to redesign the accessibility, structure and quality of psychiatric care in American nursing homes. Toward this end, this chapter will address the fundamentally important functions that may appropriately be provided by the psychiatrist in the nursing home setting.

THE ROLE OF THE PSYCHIATRIST

In the USA, contemporary training in general psychiatry emphasizes application of the biopsychosocial model to the understanding and treatment of mental disorders and behavioral symptomatology. This orientation is especially appropriate for accurately diagnosing and successfully treating the elderly nursing home resident, in whom there is often the co-morbid occurrence of physical, neurological, psychological and social contributors to disturbances of behavior. By virtue of their medical training, psychiatrists are uniquely qualified to integrate biological, psychological and social factors into a multidimensional treatment plan that reflects the full complexity of a given resident's behavioral symptomatology. Specifically, well-trained psychiatrists are potentially able to offer approaches to treatment that include recommendations for the appropriate use of a wide variety of psychoactive medications as well as psychological, behavioral

and milieu-orientated therapies. Unfortunately, little is presently known about the availability of psychiatrists to consult or manage residents in nursing homes in the USA. Additionally, a paucity of services research has been done to assist healthcare planners to better understand the character, quality and quantity of those professional functions that are provided by psychiatrists working within the nursing home milieu. In most settings, it appears that psychiatrists assume a purely consultative role, in which the primary physician (internist or family medicine practitioner) orders medication and is responsible for the course of treatment. In other settings, full responsibility for the management of the resident's psychiatric treatment resides more definitively with the psychiatrist. The factors that determine the relative intensity and scope of the psychiatrist's role vs. that of the primary physician in any given facility are largely unknown.

In a recently published study, Reichman *et al.*²⁸ examined the availability, characteristics and perceived adequacy of psychiatric consultation in nearly 900 nursing homes throughout the USA through a mailed survey to the directors of nursing of these facilities. Results indicated that 38% of nursing home residents were noted to be in need of a consultation by a psychiatrist. The frequency of these services was rated as "adequate" by only half of these homes. Nursing homes in urban and suburban regions reported better availability of psychiatric services than those located in rural areas. Nursing homes with larger bed capacities were also more likely to receive a higher frequency of services by a consulting psychiatrist. In examining the perceived adequacy of the psychiatrist's functions, these specialists were noted by two-thirds of the facilities as adequately providing diagnostic and psychopharmacologic recommendations. However, advice regarding non-medication approaches to treatment (e.g. psychotherapeutic, behavioral or milieu interventions), staff support, staff education, and attending to occasional conflict between the staff of the nursing home and resident families were reported as inadequately provided by consulting psychiatrists. Overall, the results of this study suggest that, in nursing homes in the USA, the perceived need for services provided by psychiatrists is significantly greater than the level actually provided. Additionally, it appears that the nursing directors of these settings would welcome an expanded treatment role for psychiatrists in the care of their residents with mental illness and behavioral disturbances.

While a substantial amount of research remains to be done to identify best practices for the delivery of psychiatric services in nursing homes, it is clear that the specialty of psychiatry has a vital role. In many facilities, the template of consultation-liaison psychiatry is most appropriately applied. In this framework, the psychiatrist attends to the specific mental health needs of an

individual resident, while also focusing on the needs of the nursing home staff. Resident-focused interventions provided by a psychiatrist in this context include diagnostic clarification and leadership in the assembly of a multidimensional treatment plan. For most nursing home residents suffering from mental disorders, the best outcomes likely result from optimal diagnostic accuracy and treatment that includes resident-focused psychotherapeutic or behavioral approaches, thoughtfully selected pharmacology, and specific modifications of the resident's milieu.

The liaison functions so vital to successful psychiatric care in the nursing home are varied. Clearly, the psychiatrist can be an essential resource to help staff to understand better the complex phenomenology and multifaceted treatment of dementia and mental illness in their work setting. The psychiatrist is also ably prepared to assist staff in effectively identifying and successfully managing their own job-related stress. Importantly, psychiatrists are often appropriately called upon to assist a nursing home's clinical or administrative staff in resolving conflict between the facility and a given resident's family members.

SUMMARY

In many respects, the contemporary nursing home, despite its staffing patterns and physical structure, is in large part a long-term psychiatric residential facility. Little research has been done to adequately inform the character of the best psychiatric clinical practices in this setting. However, existing data in the USA suggest that the need for formal psychiatric services in nursing homes is significant and largely under-met. By virtue of their interest and training, psychiatrists are well-suited to work as members of a multidisciplinary care team in the construction and implementation of a multidimensional treatment plan directed at a resident's mental disorder. Additionally, through the established role of liaison, the psychiatrist can do much to foster an improved sense of well-being among nursing home staff.

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Patient Autonomy vs. Duty of Care— the Old Age Psychiatrist's Dilemma

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The twentieth century has seen huge transitions in medical and legal practice based upon changes in the philosophy underpinning the way in which doctors relate to patients. At its lowest points, the Nazi programmes of extermination and forced sterilization in the USA, Sweden and elsewhere have demonstrated the capacity of doctors for abuse of their patients¹. In the UK poor-quality procedures around consent, and scandals such as patients being charged by carers for their weekly bath and other “privileges”, have shown just how easy it is for the vulnerable to be abused. Arising from this has been a philosophical, legal and medical trend towards self-determination and autonomy. Advance directives have been promoted as a solution to the loss of autonomy. They have, however, been shown to have limitations. Indeed, one study found that the entire health gain from cardiac rehabilitation programmes was neutralized because patients in the study signed advance directives².

Old age psychiatry must, if it is honest, admit that it views autonomy as a limited concept. The use of legal tools such as the UK Mental Health Act to detain mentally ill patients, along with the widespread housing of demented people behind locked doors, even without a formal detention order³, shows that autonomy does not rule undisputed. There is also evidence that the practice of administering medication covertly within foodstuffs is widespread in the UK⁴. How can we justify such acts? In a landmark judgement, the UK Law Lords held that the mentally incapacitated could be detained and treated in their best interests because

it was their illness that primarily removed their autonomy and not the fact of their detention⁵. Doctors and others may therefore treat the mentally incapacitated against their wishes when the patients themselves will clearly benefit from such treatment.

This shows that old age psychiatrists have an inescapable and awesome responsibility to balance the principles of autonomy and good clinical care when they are in opposition. This balancing act must be open to scrutiny and requires that good clinical care of the patient is the focus. The twentieth century was too heavily littered with examples of patients' needs coming second to the intentions of others for it to be otherwise. In essence, to be trustworthy, doctors can never intend harm.

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Psychiatric Services in Long-term Care

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Older people require long-term care when they have care needs that go beyond what they and their families can provide. It can consist of subacute, step-down or convalescent care for individuals discharged from hospitals, rehabilitative services for those recovering from illness or injury, and hospice care for those with terminal illness, as well as life-long care for those with irreversible disability. Increasingly, the landscape of settings for long-term care is expanding to include home- and community-based programs. Even for those who require residential care, the options are expanding to include increasingly diverse forms of assisted living or personal care facilities, as well as nursing homes. Nevertheless, nursing homes remain the most important settings for long-term care, especially for those individuals who are oldest and most disabled. However, nursing homes are also evolving, with increasing numbers of patients admitted for short stays for subacute or rehabilitation care and a proliferation of special care units designed for patients with dementia.

According to recent government reports¹⁻⁴, there are approximately 1.6 million residents occupying 1.76 million beds in 16 840 American nursing homes. For persons who turned 65 in 1990, an estimated 43% will enter a nursing home at some time. Of this group, 55% will have a total lifetime use of at least 1 year, and 21% will have total lifetime use of 5 years or more. Although both the number of facilities and the number of beds increased almost 20% from the mid-1980s to the mid-1990s, they did not keep pace with the growth of the elderly population. The ratio of nursing home beds to the size of the population aged 75 years and over dropped 17% from 1987 to 1996—127–117 beds/1000 people. However, the occupancy rate declined from 92.3% in 1987 to 88.8% in 1996. Thus, the nursing home market is beginning to experience the combined effects of healthier aging and the availability of alternative approaches to long-term care.

US nursing homes are heterogeneous: 65.5% are for-profit vs. 27.9% non-profit, 6.6% are government-owned; 53.8% are owned by chains; and 13.6% are hospital-based. In 1997, 67.4% of nursing home costs were paid by Medicaid (approximately half of this paid by the federal government and half by the states), 9.4% by Medicare, and 23.3% from private funds or other payers. Federal payments account for the majority of expenditures for nursing home care and have been estimated in the late 1990s to be approximately \$40 billion/year.

Among all US nursing home residents in 1996, the average age was 84.6; 9% were under 65, 12% 65–74, 30% 75–84 and 49% 85+; 71.6% of residents were women; 88.7% of residents were White, and 8.9% African-American; 13.9% required assistance with one or two activities of daily living tasks, and 83.3% with three or more; 88.2% required assistance with dressing, 96.5% with bathing, 59.7% with eating, 73.6% with transferring into or out of bed, 66.4% with mobility and 79.7% with toileting.

Compared to comparable findings from a decade earlier, the average age of residents increased by 0.9 years; the proportion aged 85+ increased from 49% to 56% for women and from 29% to 33% for men. Disability of residents also increased and the proportion of those requiring assistance in three or more activities of daily living was 15% higher in 1996 than 1987.

The psychiatric needs of nursing home residents are thus those of a population characterized by extreme old age and high levels of disability. Accordingly, the delivery of psychiatric services in the nursing home must be informed by knowledge of the clinical psychiatry of this population. However, mental health providers must also be aware of other factors that shape the delivery of care. These include the potential for use of the nursing home environment as a therapeutic agent, either in Special Care Units for dementia or in other programs, and the extensive federal regulations that govern clinical services in US nursing homes.

CLINICAL PSYCHIATRY IN THE NURSING HOME

According to recent reports from the Nursing Home Component of the Medical Expenditure Panel Survey (MEPS), approximately 48% of US nursing home residents have a diagnosis of a dementia¹. However, this figure probably underestimates the actual prevalence. Other MEPS data demonstrate that approximately 70% of residents have memory problems, 73% orientation problems, and 80% impairments in decision-making capacity. It also estimates that approximately 30% of residents have behavioral problems; 11.8% are verbally abusive, 9.1% are physically abusive, 14.5% are socially inappropriate, 12.5% are resistive to care, and 9.4% wander. In addition to this high prevalence of cognitive impairment, the MEPS reports that approximately 20% of residents have a diagnosis of a depressive disorder. Comparing these figures with comparable data from a decade earlier suggests that the number of residents with diagnoses of dementia or depression has increased. However, the number of individuals with schizophrenia has declined, especially among the younger residents⁵. Findings from this national representative sample confirm earlier research reports about the high prevalence of psychiatric disorders in nursing home residents⁶. In particular, they support the validity and generalizability of estimates from research in a single facility that demonstrated that 80% of residents have a psychiatric diagnosis. Findings from this earlier research provided insight into the nature of the disorders; 67% of residents had dementia, with most having Alzheimer's disease; approximately 40% of those with dementia had other psychiatric syndromes as complications (psychosis 13.5%, depression 6.3%, and delirium 7.3%); and 12.8% had other psychiatric disorders, most

commonly depression⁷. Among those with dementia, it was those with psychiatric complications who were most likely to exhibit behavioral problems.

The available findings suggest that, in spite of significant changes in American nursing homes over recent years, the high levels of psychiatric morbidity and the distribution of disorders has remained the same. There has, however, been significant progress in developing and validating treatments for psychiatric disorders in nursing homes. One particularly promising intervention for residents with dementia and behavioral problems combined augmented activities, guidelines for use of psychotropic medications, and educational rounds⁸. It was tested in a randomized clinical trial and found to be effective in reducing the prevalence of behavioral disorders and the use of both antipsychotic drugs and physical restraints. Another series of studies has shown that individualized consultations to staff nurses about the management of patients with dementia can reduce the use of physical restraints⁹. There has been little research on the effectiveness of specific psychotherapies for nursing home residents with depression. Although available findings are highly promising¹⁰, research on individualized behavioral interventions for patients with behavioral and psychological symptoms of dementia have been limited to case series or small-scale controlled studies.

Psychotherapeutic medications are widely used in nursing homes. The US Health Care Financing Administration estimated that use of these agents increased during the 1990s from 21.7% in 1991 to 46.1% in 1997^{3,4}. This reflected a 59.8% decrease in the use of antipsychotic medications from 33.7% to 16.1%, but a 97% increase in the use of antidepressants from 12.6% to 24.9%. These changes probably reflect a number of factors, including scientific developments, accumulating effects of professional education and specific federal regulations (as described below).

Two randomized clinical trials have demonstrated the efficacy of the atypical antipsychotic agent risperidone for the treatment of the psychotic and behavioral symptoms in residents with dementia^{11,12}. The available findings suggest that it has both antipsychotic effects and independent effects on aggression or agitation. Longer-term follow-up studies suggest that it may cause less tardive dyskinesia than typical neuroleptics¹³. A randomized clinical trial of olanzapine vs. placebo has demonstrated similar efficacy¹⁴ and additional findings on other atypical antipsychotic agents are expected in the near future. However, the controlled clinical trials on antipsychotic agents have evaluated only their acute effects, typically for periods of 6–12 weeks, and little is known about the effectiveness of antipsychotic drug treatment over longer periods of time. In fact, recent double-blind, placebo-controlled studies of neuroleptic discontinuation demonstrate that the majority of patients who have been receiving longer-term treatment with these agents can be withdrawn from them without ill-effects^{15,16}. Two randomized clinical trials studies evaluated the efficacy of mood-stabilizing anticonvulsants for the treatment of agitation and aggression. One studied carbamazepine and found that it was effective for agitation, hostility and aggression but not for other symptoms, such as hallucinations or delusions¹⁷. Findings from staff reports demonstrated that treatment with active medication led to decreases in the nursing time required for patient care. Another recent study evaluated valproate vs. placebo and found evidence for efficacy¹⁸.

There are now several acetylcholinesterase inhibitors approved for use in patients with mild to moderate Alzheimer's disease, and there have been questions about whether they are useful in treating nursing home residents with more advanced disease. One randomized clinical trial demonstrated that use of donepezil was associated with improvements in cognitive performance comparable to those observed in less impaired outpatients¹⁹. Although there have been suggestions that cholinesterase inhibitors may be

useful in managing behavioral symptoms, this issue is unresolved at this time.

There have been two randomized clinical trials evaluating the effects of antidepressants in nursing home residents, both using the classical tricyclic nortriptyline. One of the studies was placebo-controlled²⁰. Positive findings were used to confirm the validity of the diagnosis of major depression among nursing home residents in spite of potential confounds from medical, environmental and existential factors. Another finding from this study was that patients with major depression, low levels of serum albumin and high levels of self-care disability were less likely to respond to treatment; this led to the suggestion that patients with this clinical profile may benefit from early hospitalization and evaluation of the need for electroconvulsive therapy. The second study randomized patients to regular vs. low-dose nortriptyline and found significant plasma level response relationships in those patients who were cognitively intact, again confirming the validity of the diagnosis of depression²¹. However, the plasma level response relationship was significantly different in patients with dementia, suggesting that the depression of dementia may be a distinct disorder. There have been no randomized clinical trials of selective serotonin reuptake inhibitors (SSRIs) or related medications in nursing home residents, and available open-label studies have mixed results, especially with respect to the outcomes of treatment in nursing home residents with dementia^{22–26}. In a related area, there is evidence from an older clinical trial that the stimulant medication methylphenidate may be useful in demented patients with symptoms of apathy and withdrawal²⁷.

The recent estimate that almost 25% of US nursing home residents are receiving an antidepressant medication reflects an extraordinary change in patterns for drug utilization, especially in light of findings indicating that a generation ago only 15% of residents with a known diagnosis of depression were receiving antidepressants²⁸. Although significant components of current antidepressant usage may be for other putative indications, such as agitation, sleep or pain, it is important to note that reported utilization rates are comparable to estimates for the prevalence of depression. Research is needed to determine whether it is possible to demonstrate an impact of prescribing on the mental health of the population as a whole.

THE NURSING HOME AS A MENTAL HEALTH CARE ENVIRONMENT

In the past decade there has been a growing awareness of the importance of the psychosocial and mental health aspects of nursing home care and a number of conceptual models have been developed to focus on the person–environment interaction as the target for care practices. These have included “progressively lowered stress threshold”²⁹, “stimulation-retreat”³⁰, and “person-centered”^{31–33} care. Other widely discussed concerns that reflect the increased awareness of the psychosocial aspects of nursing home care include quality of life³⁴, individualization of care³⁵, the importance of the patient's perspective^{36,37}, and autonomy^{38,39}.

Many of these concepts have been applied in the design and operation of special care units (SCUs) for residents with cognitive impairment. The popularity of these units has been phenomenal, with current estimates that 22% of nursing homes have designated SCUs for patients with dementia. Outcome studies evaluating special vs. traditional care have suggested positive effects in a variety of selected outcomes, such as the nature of the services provided, depression, family perception of quality of life and the rate of decline in mobility^{40–43}. However, more rigorous randomized clinical trials have reported positive outcomes only in circumscribed areas, such as catastrophic reactions⁴⁴ and observed positive emotional responses⁴⁵.

While it is encouraging to find evidence that the outcomes of care in nursing homes may be modifiable, it is important to recognize that there are no effects that are consistent across studies. However, these mixed findings are not unexpected, given the tremendous variability in the definitions, structure and programs of SCUs as well as other methodological limitations, such as selection biases, attrition and measurement limitations. Therefore, many researchers have increasingly called for studies focused on the evaluation of specific interventions associated with improved comfort, health status and quality of life, in attempts to identify and evaluate the “active ingredients” of special care^{37,46,47}.

Many programs have been based upon modification of the physical environment to decrease behavioral difficulties and enhance positive quality of life in the nursing home. Promising programs that incorporate natural elements into the environment, include a bathing intervention that uses bird songs, pictures of nature and food during bathing to decrease agitated or aggressive behavior⁴⁸, and use of “white noise” machines emitting sounds of ocean waves or waterfalls⁴⁹. The use of bright lights in the care environment has been shown to consolidate sleep and to reduce agitation, but only in those with disturbed sleep-wake cycles⁵⁰. An intervention that decreased noise and light in the night-time environment did not, by itself, improve the quality of the residents’ night-time sleep; however, when combined with a daytime program of increased physical activity, it did provide benefits^{51,52}.

Some programs have gained great popularity in the clinical community, but have not yet been evaluated in controlled studies. These include the “Eden Alternative”, which attempts to increase biological diversity in the institutional setting by incorporating children, plants and animals into the day-to-day life of the nursing home⁵³, and horticultural interventions⁵⁴. Another set of promising interventions are based upon spirituality or religiously-based programming^{55,56}. Music therapies, in particular, have been studied more intensively and may be effective in increasing positive emotional and social engagement and decreasing problem behaviors⁵⁷.

Technological interventions have been used in the nursing home, either to provide enhanced surveillance to increase safety or to use modalities such as audio or videotapes as therapeutic tools. Only one of these therapeutic interventions has shown positive results after testing under controlled conditions—simulated presence therapy⁵⁸, in which residents with dementia are asked to listen to taped, simulated telephone conversations with family members, was superior to placebo in decreasing problem behaviors and increasing well-being.

Several studies have demonstrated efficacy in programs designed to promote functional independence in persons with dementia residing in the nursing home. One focused on altering “dependency support scripts” and found increased independent behavior on the part of residents⁵⁹. Other studies have focused on evaluating programs designed to target specific functional capabilities. Several groups have found positive effects from programs that modify usual routines for morning care and dressing. One analyzed videotapes of dressing behaviors to develop care prescriptions, and found that these led to increased independence in dressing⁶⁰. Another used a “skill elicitation” intervention and found that it significantly increased the amount of time participants were engaged in dressing and other ADLs, while simultaneously decreasing the frequency of disruptive behavior⁶¹. A third found that an “abilities-focused” program, teaching direct care staff to modify the moment-to-moment procedures used in morning care on the basis of knowledge of the residents’ specific cognitive deficits, led to improvement in calm/functional behaviors, agitation and social functioning⁶².

Other important factors affecting the psychosocial environment in nursing homes are those related to the caregiving staff. It is

certified nursing assistants (CNAs) who serve as primary caregivers to a population characterized by ever-increasing levels of dementia, dependency and mental health disorders. There is no doubt that the increased demands on skills and time, together with the physically and emotionally demanding labor, require attention to selection, training, supervision, and a focus on the role of the CNA as a provider of psychosocial care. There are multiple sources of stress for these paraprofessionals, including individual self-related needs, off-the-job stressors, patient contact, and administrative and organizational factors. One study demonstrated that 57% of CNAs screen positive for clinically significant levels of distress⁶³. However, the paraprofessional staff do not, as a rule, receive adequate support from the psychological and psychiatric community in addressing such issues as non-pharmacologic management techniques, assisting with family conflicts and coping with job-related stress⁶⁴.

Fortunately, there is evidence that training programs for CNAs can have positive outcomes for the CNAs, including increased interaction and sensitivity to resident cues, provision of more choice and praise, and increased behavioral management skills. Moreover, these can translate into improvements in residents’ mood and functioning^{65,66}. Another line of research has found that mental health outcomes can be affected by structural and organizational elements of nursing homes, such as chain status, size, staffing levels, turnover, staff selection, job assignments, job design and the adequacy of supplies; moreover, interventions modifying formal elements of staff management can have positive effects on staff behavior⁶⁷.

THE REGULATORY ENVIRONMENT IN US NURSING HOMES

Although nursing homes have historically been designed to care for patients with medical and surgical conditions, the vast majority of their residents have psychiatric disorders. The mismatch between resident needs and facility characteristics in US nursing homes has been associated with inadequate, inappropriate and even inhumane treatment⁶⁸. During the 1980s, concerns expressed by advocacy and professional groups were reinforced by a report from the Institute of Medicine⁶⁹ documenting major problems in the quality of care provided in nursing homes. Specific issues included the undertreatment of depression and use of physical and chemical restraints to control behavioral symptoms in patients with dementia. Other concerns at that time were that elderly patients with chronic and severe psychiatric conditions were discharged from state hospitals and inappropriately placed in nursing homes at Medicaid expense, thereby denying them access to the active psychiatric treatment they needed, and shifting a substantial portion of the costs of their care from the states to the federal government.

Recognition of problems in the quality of care in US nursing homes, together with ongoing concerns about costs, prompted Congress to pass legislation, the Nursing Home Reform provisions of the Omnibus Budget Reconciliation Act (OBRA) of 1987⁷⁰, which has transformed nursing homes into one of the most highly regulated environments for healthcare delivery in the USA. To operationalize the laws enacted under OBRA ’87, Congress directed the Health Care Financing Administration (HCFA), the agency that administers Medicare and Medicaid, to issue specific regulations that govern nearly all aspects of nursing home operations⁷¹, and charged the states with the responsibility for conducting surveys to determine whether nursing facilities were in compliance. In response, HCFA developed a set of interpretive guidelines for state surveyors⁷², which has been revised over the past 10 years to reflect changes in healthcare practice and policy. Mental health screening, evaluation, care

planning and treatment are addressed under sections of the regulations related to resident assessment, resident rights, facility practices and quality of care.

The OBRA regulations require preadmission screening and annual resident review (PASARR) to prevent inappropriate nursing home admissions for patients with severe psychiatric disorders, and to ensure that those with acute psychiatric conditions are not placed in nursing homes before receiving the benefits of acute psychiatric treatment⁷³. However, patients who have a primary diagnosis of dementia are considered eligible for nursing home admission regardless of psychiatric complications or co-morbid conditions. After admission to a nursing facility, all patients are required to undergo periodic comprehensive assessments using the Minimum Data Set (MDS), a standardized instrument that includes several areas relevant to mental health: mood, cognition, communication, behavior patterns, activities, functional status, psychosocial well-being, oral/nutritional status, co-morbid disease, medications and other treatments⁷⁴. Responses on the MDS that indicate deficits or changes in the patient's health status serve as triggers for resident assessment protocols (RAPs), which are second-stage assessment tools that furnish prompts to help nursing staff recognize signs and symptoms that are indicators of potentially significant clinical problems, algorithms to direct further evaluations, and guidelines for treatment planning. RAP problem areas related to mental disorders and behavioral health include delirium, cognitive loss/dementia, psychosocial well-being, mood state, behavior problems, psychotropic drug use and physical restraints. The regulations hold facilities responsible for ensuring that the MDS is completed on time and the RAPs are followed.

The OBRA regulations also contain provisions that affect psychiatric treatment in the nursing home, prohibiting the use of physical restraints or psychotropic drugs when they are "administered for purposes of discipline or convenience and not required to treat the resident's medical symptoms" or promote improved functioning⁷¹. There are specific regulations concerning antipsychotic drugs, designed to ensure that residents receive these agents only when "necessary to treat a specific condition as diagnosed and documented in the clinical record". The regulations also require periodic "gradual dose reductions and behavioral interventions, unless clinically contraindicated, in an effort to discontinue these drugs". Syndromes such as delirium and dementia, when accompanied by agitated or psychotic features, are listed among the accepted indications for the use of these agents. However, use of these medications must be supported by documentation that there are specific target symptoms associated with resident distress, functional impairment or danger to self or others.

Regulations related to quality of care further require that each resident's drug regimen be free from unnecessary drugs, defined as any drug used in excessive dose (including duplicate therapy); for excessive duration; without adequate monitoring; without adequate indications for its use; or in the presence of adverse consequences indicating the need for dose reduction or discontinuation⁷¹. The interpretive guidelines that accompany these regulations specifically limit the use of antipsychotic drugs, antianxiety agents and sedative-hypnotics, but not antidepressants⁷². For each of these medications, the guidelines provide a list of acceptable indications as well as specific agents that may not be used; daily dose limits which, if exceeded, require documentation that the benefits justify the risks; requirements for monitoring treatment and adverse effects; and time frames for attempting dose reductions and discontinuation. These guidelines were updated in 1999 to reflect new clinical knowledge and accepted practice and to include newly approved drugs.

Beginning in July, 1999, HCFA introduced quality indicators (QIs) derived from the MDS to enable surveyors to compare

individual facilities to others in the same state⁷⁵. There are 24 QIs within 11 different domains, including behavioral/emotional problems, cognitive patterns and psychotropic drug use. Behavioral/emotional patterns cover the prevalence of behavioral symptoms affecting others (e.g. verbally or physically abusive, socially inappropriate or disruptive behavior); prevalence of symptoms of depression; and prevalence of depression without antidepressant therapy. The cognitive pattern domain examines the incidence of cognitive impairment when consecutive MDS assessments reveal new onset of impairments in short-term memory or decision-making ability. The psychotropic drug use domain includes the prevalence of antipsychotic use for patients without psychotic conditions, the prevalence of antianxiety/hypnotic use, and the prevalence of hypnotic use more than twice in the previous 7 days. QIs that are indirectly related to mental disorders and their treatment include the use of nine or more different medications, and the prevalence of falls, weight loss, daily physical restraints, and little or no activity. Whenever a review in any of these areas results in a deficiency citation, a plan of correction must be developed and approved.

SUMMARY

In spite of significant changes in American nursing homes over the past 10–15 years, the high prevalence rates for dementia and depression remain as persistent problems. Fortunately, there have been major advances in knowledge about the outcomes of psychiatric care for nursing home residents. Chapters in textbooks from a decade ago may have been eloquent about the need for mental health services, but they had little to say about their outcomes. Now, it is possible to review evidence for benefits of specific interventions. In addition to pharmacological and behavioral treatments that are designed to be delivered to individual patients, other interventions are designed for delivery to nursing units or to residential care facilities as a whole. Promising interventions in this domain include augmented activities, modifications of the physical environment, use of technologies to augment interpersonal interventions, modifications in patterns for the delivery of basic nursing care, staff education, and changes in administrative structures. All of these must be delivered in the context of a complex regulatory environment. Thus, the delivery of psychiatric services in nursing homes must attend to the needs of the individual patients and to the context in which they receive care. To be successful, it requires partnerships between mental health professionals, direct care staff and the facilities' administration.

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Care in Private Psychiatric Hospitals

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Psychiatric care and treatment for mental disorders in the elderly in a private psychiatric setting is fundamentally a public health issue. The location of treatment and the clinical services available are shaped by public policy, financial incentives and the emergence of managed care. With the implementation of the Medicare Part A Hospital Insurance Trust Fund, most acute inpatient psychiatric care and treatment for the elderly in the 1970s and 1980s occurred in special geropsychiatric units in general medical units and private psychiatric hospitals. Data reflect the trend of the shift from public sector inpatient care to private and general medical hospital inpatient units. State and county organizations providing mental health care decreased 7.6%, while private hospitals increased 95.5% and general medical hospitals with inpatient mental health services increased 21.6%. The relative percentage of mental health expenditures for state and county hospitals decreased from 48.5% in 1975 to 23.6% in 1994 for total mental health expenditures; while the percentage of expenditures for private psychiatric hospitals increased from 7.1% to 19.5% during that same time period. Outpatient and community-based programs, including partial hospital programs, increased as a relative percentage of total mental health expenditures from 1.8% in 1975 to 26.8% in 1994¹. In addition to addressing concerns inherent to providing quality medical care, any discussion focused on the care of older persons must include content regarding the logistical and ethical dimensions of financial and familial responsibilities. In the context of an increasingly aged population, Henderson² challenges us with a public health agenda of: (a) developing "bold and innovative means for assisting families to care for a relative with dementia"; (b) improving the "contributions of general practice to the care of mental disorders in the elderly"; and (c) investigating the social environment of the mentally ill elderly and promoting more adaptive alternatives when feasible. These challenges are particularly pertinent in the private psychiatric facility, where the "financial disincentives and shortages of trained personnel"³ common to the discipline of geropsychiatry are frequently amplified.

There are several factors that necessitate the private psychiatric community becoming more aggressive and innovative in the care of older persons, despite the substantial constraints and disincentives. The first is the void left by the inadequacies in the community mental health center and the shift from treatment in state and county hospital programs for this population⁴. Second, the overall demand for inpatient psychiatric treatment for the elderly has continued to grow as the number of elderly, especially those over 80, has grown dramatically⁵. A third factor leading to an increased demand for private psychiatric care is that internists

and primary care physicians are frequently not trained in the behavioral management of geriatric patients, and there is a growing subspecialty of geriatric psychiatry which is generating paradigms for diagnosis and treatment of older persons with neuropsychiatric and behavioral disorders.

The World Health Organization has identified acute inpatient psychiatric care as an important component in the continuum of care for the elderly⁶. Although the USA and the American Psychiatric Association have not published specific practice guidelines for geriatric inpatient treatment, there are generally accepted principles of quality care and treatment of elderly individuals in inpatient settings. These principles of care include: preadmission screening and linkages with community providers of care; comprehensive assessment and care planning; multi-disciplinary staff with specialized experience and interest in the elderly; ongoing staff training and education; therapeutic programming and care approaches sensitive to the needs of an aging population; environmental and physical design of program; individual, group and family therapy; discharge and aftercare planning. With the emergence of managed care and financial incentives for community care, there has been a lack of coordination and integration of mental health services, especially for the elderly with severe mood disorders, psychosis, and dementing illnesses with complicating psychiatric and behavioral disorders. These forces combine to magnify significant gaps in treatment for the most frail and psychiatrically needy elderly who need comprehensive psychiatric and medical treatment, most appropriately provided in an acute inpatient geropsychiatric treatment program. Keill⁷ proposes that it is "still possible within the system and with proper incentives to provide an accessible, comprehensive network". He emphasizes that within this network there must be a continuity of care, which is defined by Bachrach⁸ as "a process involving the orderly, uninterrupted movement of patients among the diverse elements of the service delivery system". In the context of these challenges and stipulations, we present an example of a geropsychiatry service in a private hospital through which such issues can be discussed.

DESCRIPTION OF PROGRAM

The Parthenon Pavilion at Centennial Medical Center in Nashville, Tennessee, is a 162 bed private proprietary psychiatric hospital that has developed a cost-effective model program for the delivery of acute inpatient geropsychiatric services. The hospital currently operates two 12 bed Alzheimer's disease and related disorders-memory disorders units and a 16 bed general

geropsychiatry unit, which together function as a regional referral center for the treatment of the elderly with complex and concurrent psychiatric and medical problems. Parthenon Pavilion's geriatrics program opened in 1985 and the original memory disorders unit was developed in 1987. This program provides a structure for supporting continuity of care by organizing clinical services and concentrating resources around the diverse and specialized needs of patients with severe psychiatric disorders, often complicated by behavioral difficulties and/or serious medical problems. The hospital also includes a geriatric partial hospitalization program which adds another dimension to the continuum of psychiatric care for senior adults.

Clinical Process

The geriatrics program has developed a philosophy of treatment and a clinical services model that places an emphasis on interdisciplinary care, so as to provide psychiatric and medical management of treatable symptoms in order to reduce distress, disability and complications in the older patient. A second major tenet of the program's treatment philosophy is that the family and professional caregiver systems are helped by providing: (a) accurate diagnosis and education around the nature of disease processes in the context of the unique manifestations of a disease in a given individual; and (b) supportive assistance and practical guidance tailored to the care and management issues of a given patient and caregiver system. This clinical services model also places a high priority on providing consultative and clinical liaison services to families, agencies, nursing homes and residential facilities, to assure continuity of care and maintenance of the clinical goals established during hospitalization for patients after they are discharged from the program.

The importance of family involvement in such a program cannot be overstated. Hardwig⁹ challenges the ethically simplistic notions of patient autonomy frequently espoused when he points out the integral role of the family in making medical decisions. This is especially relevant when working with older persons, due to the prevalence of cognitive impairment found in this group. When a patient becomes demented, family members must frequently assume decision making in medical situations. This may violate family taboos as well as established patterns of family interaction, which may require renegotiation. Family conferences, as a means for mediating the moral process of medical decision-making and planning for ongoing care, are an integral aspect of the clinical services offered by the Geriatric Program, and result in a consistently high level of family satisfaction.

Administrative and Team Function

The administrative structure of this program includes: the clinical director, who is a psychiatrist specializing in geropsychiatry; a program director who is a Master's-trained social worker; a nurse manager with experience in geriatric nursing; and access to a clinical nurse specialist with a Master's degree in psychiatric nursing. The clinical director provides leadership in program planning, program evaluation and quality assurance, while also being available to provide consultation to other attending psychiatrists as needed. The program director coordinates the day-to-day operation of the program and works closely with the clinical director and nurse manager in implementing the above-designated functions.

Each patient has an attending psychiatrist, who directs the treatment, meets with families in diagnostic feedback conferences, participates in clinical liaison activities with nursing homes and

other placement facilities and develops an aftercare plan, along with other disciplines represented on the staff¹⁰. The program is supported by a number of psychiatrists with added qualifications in geriatric psychiatry and considerable experience in the treatment of such patients. Although many private psychiatric facilities have limited professional staff with particular expertise in geriatrics, such persons are increasingly available, due to an increasing number of specialized training programs. The program is joined by a group of board-certified internists, who have special interest and expertise in evaluating and treating acute and chronic medical problems in the elderly. Upon admission, the internist conducts a comprehensive review of systems and physical examination on each patient. Throughout the hospitalization, the internist provides follow-up of existing or developing medical problems and, when necessary, participates in family conferences and team staffing. Medical subspecialists, such as neurologists and cardiologists, are consulted when clinically indicated and their availability is fostered by the fact that the hospital/program is part of a regional tertiary care medical center which also provides access to available technology, brain imaging, laboratory services and specialized neuropsychological testing as medically indicated.

Nursing care is delivered through a primary nursing system. Nurses are recruited who have strong medical/surgical background along with geropsychiatric interests¹¹. Meeting these staffing demands remains challenging, particularly in an era when most hospitals are experiencing professional nursing shortages. Supportive nursing administration within the program and the promotion of educational opportunities have fostered a reduced staff turnover rate well below the 20% annual rate experienced nationally¹². The program director and the nurse manager are involved in pre-admission screening and clinical liaison activities, along with providing inpatient clinical services to family and professional caregivers. The social workers coordinate discharge planning and aftercare follow-up, which is a vital linkage in maintaining continuity of care. Activity therapists conduct functional assessments and design appropriate activities for the units within the program, according to the level of function of the patients. They can provide feedback to families and professional caregivers regarding functional abilities and deficits, making recommendations for modifications in the patient's environment and activities to support remaining abilities. Staff education is conducted through a variety of means, including weekly teaching rounds conducted by the clinical director, monthly interdisciplinary staff education meetings, and special teaching modules designed by the clinical nurse specialist.

Physical and Environmental Features

The physical design of the geriatric program provides for separate units divided by a nursing station. Two are for patients with dementia and memory disorders; the other for the general geriatric population. Prior to the development of the specialty units, demented patients were integrated with non-demented patients and separate therapeutic activities and groups were planned for each population¹³. However, it ultimately was determined that separating the program into specialty units, serving cognitively impaired patients with Alzheimer's disease and related disorders and a general geropsychiatric unit provided the optimal arrangement both clinically and administratively. The memory disorders units are designed for security and have unit doors that can be locked. Such a unit is specifically designed and adapted for the particular needs of sensory and cognitively impaired persons. There is an emphasis on music, videotapes and appropriate sensory stimulation without overload. The general geropsychiatry unit and the memory disorders units are pod-shaped, with a day room outside the bedrooms, and have activity

rooms that can be used for special activities as needed. The care environments are designed to be adaptive in nature and focus upon remaining rather than lost abilities.

Financial Considerations

A necessity in the private psychiatric hospital is to operate in a cost-effective manner, so that the program does not have a significantly negative financial impact on the hospital. One means of achieving this goal has been to establish a program-based preadmission screening and consultative service to assure that each person admitted to the program meets Medicare intensity and severity criteria for a psychiatric admission. The pre-admission service also screens for persons primarily needing medical treatment or long-term care, and focuses on orientating the family and referral sources to the goals and limitations of inpatient psychiatric treatment. Consultative and crisis management assistance is provided when hospital treatment is either inappropriate or not immediately available.

Another means of maintaining cost-effective utilization of resources is to avoid inappropriately extended stays by assuring that discharge planning begins with the preadmission process and is an integral aspect of ongoing treatment. Length of patient stay is monitored through a case review-orientated quality assurance program, coordinated by the program director working in cooperation with the clinical director. Regular review of the length of stay and systematic documentation of reasons for continued hospitalization not only enhances the quality of care but also provides a peer-reviewed justification for continued stay when reviewed by regulatory agencies. A major challenge during the past decade has been the shift to Medicare Managed Care, an HMO, for cost containment and utilization. A goal of such programs is to achieve more cost-effective care and treatment and to prevent unnecessary hospitalization. However, aggressive cost containment can negatively affect access and quality of care for the elderly with severe mental disorders and dementia, complicated by co-morbid medical conditions. This effort has resulted in undertreatment and limited access to inpatient care. In addition, with psychiatric care "carved out" of medical benefits and financial incentives to limit care, the distinction between medical and psychiatric coverage in demented patients may have significant financial impact on third-party payers.

Summary

In summary, the key elements of the program used in the model emphasize: (a) preadmission assessment and screening in the context of consultative services; (b) treatment by an interdisciplinary team, which includes an attending psychiatrist, internist, primary nurse, social worker, activities therapist and other ancillary staff, each with a commitment to the particular needs of the older person; (c) individualized educational programs for families and professional caregivers to augment existing clinical services; (d) individualized behavioral approaches that incorporate written behavioral management plans as an adjunct or substitute for psychopharmacological approaches; (e) discharge and aftercare planning services, family conferences and post-hospitalization follow-up with families and institutional caregivers; (f) environmental and physical design; and (g) cost-effective inpatient treatment. Approaches for assuring continuity of care for individual patients with complicated behavioral and psychiatric symptoms have included written behavioral prescriptions for families and caregivers, demonstrations of appropriate caregiving techniques during hospitalization, and videotape demonstrations designed to serve as training tools for nursing

home staff members. In 1999, Parthenon Pavilion provided leadership and financing for a new initiative to focus on improved mental health care for older adults. The initiative represents an innovative approach to clinical collaboration among select organizations providing mental health services to the elderly. The clinical Senior Links mental health consortium included representatives from home health, the local mental health association and Alzheimer's association, a community mental center and senior center providing community-based mental health services, staff of Parthenon Pavilion, and geriatric partial hospital programs located in the community. This program serves as a potentially innovative approach to improve the delivery of mental health services for senior adults and provides a forum for focusing on improved access, assessment and treatment, aftercare coordination and education for the consortium members and the community.

CHALLENGES FOR THE FUTURE

The primary challenges facing private psychiatric hospitals serving the elderly include: (a) monitoring and modifying admission criteria as Medicare intensity and severity criteria continue to become more restrictive and prospective payment is implemented; (b) developing intervention strategies with family caregivers to avoid inappropriately extended stays for non-clinical reasons; (c) maintaining quality care in the context of financial disincentives, especially magnified by increased enrollment in Medicare managed care plans, and regulatory policies which impact negatively on hospitals serving geriatric patients with severe psychiatric illnesses frequently complicated by behavioral and medical problems; (d) maintaining quality staff in all disciplines, especially with the growing nursing shortages and shrinking revenues; (e) developing consultative and outreach services to respond to the growing trend for nursing homes to serve increased numbers of residents with mental disorders and severe dementia with psychiatric and behavioral disorders.

We face these challenges in a cultural and federal regulatory environment that is ambivalent at best in its support of Medicare. Medicare accounted for only 9% of the federal budget in the USA in 1989, but sustained 36% of all budgeting cutbacks. Of even greater consequence for the elderly with mental disorders is the fact that Medicare spends only 3% of its budget on mental health care¹⁴. The claims are made that "key business values, including attention to a guiding mission statement, the needs of consumers, accountability and marketing, have a positive impact on the quality of milieu treatment"¹⁵, but these assertions are put forth in a consumer-market paradigm and the pitfalls of this approach have not been adequately assessed. Tischler¹⁶ points out the lack of systematic studies that include qualitative intermediate or long-term outcomes of mental health care. Studies examining utilization review outcomes have included only service utilization and expenditures. The program model presented here is committed to perpetual quality improvement, but clinical quality and reimbursement issues are inextricably linked in the private setting and future excellence may be jeopardized if current trends continue.

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Quality of Care and Quality of Life in Institutions for the Aged

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Although the form and function of the institution for older people changed considerably over the second half of the twentieth century in the USA, UK and other countries, the central task endures—providing protective care for older people unable to care independently for themselves. At the beginning of this period there were still people in the system whose lack of independence was primarily financial. Physical ill-health and, later and into the present, mental and cognitive ill-health became the main reason for entry into an institution.

The details of how such care was delivered differed in major ways across countries, to the point where knowledgeable comparisons were possible only for highly specialized experts who had the time to become familiar with more than one system. The present chapter is written when such differences are still evident. It is thus necessary at the outset to acknowledge that any attempted generalization about institutions for the aged must be interpreted in the light of differing cross-national social and cultural traditions. Nonetheless, this chapter will assert that there are characteristics common across localities that are universally accepted as indicators of quality of care and quality of life in institutions for older people (sometimes referred to as “nursing homes”, as in the USA).

After establishing the importance of the quality concept, this chapter will review issues in maintaining quality in this type of residential care. Some of the recent literature will be reviewed, followed by the presentation of a model for defining quality of care and quality of life now under development by Rosalie Kane, Robert Kane and the author.

WHY BE CONCERNED ABOUT QUALITY IN NURSING HOMES?

Quality has been an issue among US nursing homes from the beginning of such institutions. Governmental monitoring of quality has been a threat to for-profit entities because their profits might be threatened by external quality-monitoring efforts. Poor quality may also be found among governmentally sponsored and non-profit homes. Suffice it to say that there has always been a major gap between accepted standards and the actuality of care, sometimes to a shockingly unacceptable degree, in the USA.

Governmental regulation, although resisted by many elements of the nursing home network, has been the major device used to enhance the quality of nursing home care in the USA. The system now in place requires each state to hire and train professionals

(“surveyors”) to spend time at least every second year on site with staff, residents and archival records to assess each institution on a series of written standards.

The procedure for monitoring nursing home quality required by the US government is instructive in defining current standards. There are 185 such regulations, which are organized into 15 categories: resident rights; admission/discharge rights; resident behavior and institutional practices; quality of life; resident assessment; quality of care; nursing services; dietary services; physician services; rehabilitation services; dental services; pharmacy services; infection control; physical environment; and administration¹. Classification problems are immediately apparent. All categories are aspects of quality of care, and many may also reflect quality of life. Both of these subcategories are defined for monitoring purposes in a much more limited way than seems appropriate to this author. In 1997 the 10 most prevalent deficiencies in US nursing homes were: food sanitation; resident assessment; care plans; accidents; pressure sores; quality of care; restraint use; housekeeping quality; dignity; and accident prevention².

Citation for deficiencies may result in fines, temporary suspension of reimbursement for care or, at worst, removal of a license and closing of the institution. Suspension of reimbursement or license rarely occurs; in fact, the problems of finding care for residents in the offending institution are viewed as more stressful than continued low-grade care. Regulation is quite different, of course, in the UK. For instance, until recently it was only privately-administered facilities that were subjected to outside regulation, on the theory that local authorities were, by definition, assuring adequate quality by the nature of their direct responsibility for care³. This separation has changed, however. It seems likely that the phenomena of diversification of sponsorship (especially into the for-profit sector) and the need for local authorities to lean more heavily on professionals for quality controls than on local monitoring will make the two countries' systems become more similar in the future.

DEFINING QUALITY OF CARE

The classic system view of the health-care institution denotes the structural, given characteristics as input, care and treatment as process, and the resultant effect on the patient as output⁴. It has been noted frequently that output is difficult to identify when the institution constitutes the last residence and the final outcome is death. In addition, research over the past couple of decades has

demonstrated the dynamic character of the structure of institutions. Administrative, care-delivery and physical environmental changes in fashion, including remodeling and simple space-use changes designed to encourage desired behaviors, attest to the ability of most institutional elements to be shaped toward higher-quality care⁵.

Thus the literature of quality of care has been characterized either by value-based assertions that particular processes were intrinsically associated with higher quality of care, or were statistically correlated with overall quality as assessed by experts. The Institute of Medicine (IOM) report on nursing home quality called for making direct assessments of performance outcomes in terms of clearly undesirable states, such as death rate, infection rate, decubitus rate, or malnutrition⁶. More recently, other indicators reported on the required periodic assessment contained in the Minimum Data Set of the US nursing home system (MDS) have included accidents, questionable medication use, restraint use, infections and other obviously undesirable outcomes^{7,8}.

DEFINING QUALITY OF LIFE

Clearly, poor-quality care will lower overall quality of life. A distinction between quality of care and quality of life is useful to make, however. In a rough sense, quality of life must include features of everyday life that enhance enjoyment and sense of hope or purpose above the average level. An individual's quality of life is a subjective assessment made by that person alone. Quality of life of an environment, such as a nursing home, is represented by institutional attributes that have a statistical probability of leading to higher individually perceived quality of life for its occupants. There is thus a dual perspective on quality of life, the individual-subjective and the environmental-objective⁹.

If quality of life is to be monitored, both perspectives must be assessed. Ideally, positive features of residential care would be identified by their association with positive subjective responses by a majority of the consumers of such care. The importance of the consumer was recognized in the regulations that followed passage of the Nursing Home Reform section of 1987 legislation¹⁰. Nursing homes are required to solicit opinions on the quality of care and quality of life experienced by residents.

In practice, however, it is not possible to demonstrate a direct parallel between nursing home resident perspectives and features that represent quality in the institution. Until the state of the art of measuring both consumer evaluation and environmental attributes is further advanced, it is necessary to make many assumptions about what constitutes quality, based on the available literature. Thus, the assessment system must encompass a large array of both personal and environmental features.

A Conceptual Basis for the Search for Quality

Kane, Kane and Lawton¹¹ have found it convenient to organize quality into 11 domains: security; functional competence; comfort; dignity; autonomy; privacy; meaningful activity; social relationships; enjoyment; individuality; and spiritual well-being. These domains represent universal individual needs, whose satisfaction may be enhanced or blocked by the environment in which the person pursues the gratification of needs. In overview, an approach to assessment evaluates the extent to which residents' needs are fulfilled and the extent to which environmental features relevant to these needs are present. Although the actual design of the measures is still in process, their components may be viewed as a model that could be useful for later investigators.

Resident Needs

Many modes of consumer assessment have recently become available^{12,13}. Our own approach queries residents systematically about their evaluation of how well the residential environment fits each need. All such direct approaches require ordinary comprehension of questions and the willingness to respond frankly. Because many people in residential care are cognitively impaired and others may be loath to express critical comments, other complementary or parallel sources of information must be sought.

Resident Needs as Perceived by Others

Caregiving staff and family members are an obvious source of information on some domains. Such characteristics as functional health, cognitive performance, participation in activities or depression may be rated by an outsider. Some intrinsically subjective domains are less amenable to these types of judgments, e.g. the degree to which dignity is experienced in nursing home life. An outsider may assess a resident's ongoing affect states but clearly is limited in access to the resident's actual happiness, sadness or other feeling states¹⁴.

Direct Observation

On the other hand, systematic observation by research staff or quality-control staff may reveal very concrete instances of behavior relevant to quality of care and quality of life. An observer may be trained to be less susceptible to bias than the resident in terms of denying socially unacceptable behavior, and is capable of being instructed in the subtle indicators of quality that may be exhibited in settings such as morning care, mealtimes, activities or unprogrammed time. Examples from earlier research include systematic observation of the "behavior stream" or the non-verbal indicators of emotional state^{14,15}. It is also possible to train experts in more global aspects of direct observation that focus on concepts rather than small behavioral acts. The Professional Environmental Assessment Protocol (PEAP)¹⁶, for example, requires an environmentally trained professional to spend about an hour in a care area, after which global ratings are made on the environment's ability to foster orientation, safety and security, privacy, stimulation quality, regulation of stimulation, functional competence, personal control, and continuity of self¹⁷. One important way in which direct observation adds to the quality attainment process is that it allows for the input of expertise in judgments of quality. Not all goal-relevant information is evident to the consumer. Some of what is learned from observation is thus complementary to the resident's perspective.

Integrating the perspectives of residents, significant others and objectively-viewed phenomena is not a straightforward process. Although most experts would wish to give primacy to the views of residents themselves, around 20–40% are cognitively unable to express evaluations and preferences that might guide the enhancement of quality¹⁸. It might be argued that the most-capable 60% should be able to articulate a consumers' view that would also fit the cognitively impaired. We must recognize, however, that major impairments in cognitive and self-care ability may also translate into needs quite different from those who are intact. The perspectives of significant others and value-judgments based on observable behavior clearly add something to knowledge about quality. Yet we cannot automatically substitute them for the absent judgments of those who are too impaired to be questioned. At best, putting together the three perspectives is at present more an artistic endeavor than a scientific one. How one accounts for biases in perspectives or for the differential weighting

of the several types of input into measuring quality are tasks for the future.

The Economic Perspective

This chapter cannot do justice to the economic ramifications of quality of care and quality of life. In general, increased quality comes at a cost. In the case of both the profit-seeking institution and the governmental or non-profit institution, there are obvious constraints on the extent to which higher quality can be financed. This observation helps identify another distinction between quality of care and quality of life. Quality of care, being a life-and-health issue, is capable of being defined in terms of minimally acceptable threshold levels. These levels can in turn be defined reasonably clearly and used as the basis for licensing or potential decertification.

CONCLUSION

It is a different issue whether what has been defined as quality of life can be audited and used legally to improve the quality of nursing homes. Surveyors could conceivably become equipped to diagnose quality-of-life deficiencies and cite them to the point of removal of licensure. This appears problematic because quality of life has so many subjective aspects. Surveyors would no doubt be reluctant to make such citations and political pressure from owners would minimize the legal clout of such citations. One possible outcome is that the present survey process will continue to be used to correct the most egregious lapses in minimum quality of care. Beyond the merely adequate quality of care attained by correcting basic deficiencies, improvement in quality of life up through the positive to excellent ranges may be a matter better controlled by the marketplace than by legal enforcement. Quality-of-life audits could lead toward intrafacility self-assessment, staff training and growth, articulating possible avenues for improvement on which administration and staff could work proactively. The greatest weakness of the marketplace hypothesis, however, is that the present market is responsible primarily to the upper income range of client families. If market-driven improvement in quality of life is to occur, we should have to see greater equalization of opportunity and increasing competition for the patronage of government-subsidized residents as well as those who pay in full for their care. Despite such difficulties, we should leave room for the possibility that improved quality of life in nursing homes may emerge better with indigenous rather than legal motivation. In the UK, there is some hope that the cultural and ethical norms may support the general public and governmental view that quality of care and quality of life are both basic rights of all citizens. This motivation might over time become more effective than either legal regulation or market competition¹⁹.

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Liaison with Medical and Surgical Teams

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In one of the early textbooks of geriatric medicine, Agate¹ wrote, "As a sign of acute physical illness in old age, mental change is more significant than a rise in temperature or pulse rate...". Not only is delirium (acute organic brain syndrome) more frequently found in old age than in earlier life, but the incidence of dementia rises sharply and affective disorders, particularly depression, remain common. Thus, someone presenting with medical or surgical problems may well have coincidental, as well as causal or resultant, psychiatric symptoms. It is hard, therefore, for physicians and surgeons to ignore the psychiatric problems of their patients.

PREVALENCE OF PSYCHIATRIC DISORDER IN NON-PSYCHIATRIC INPATIENTS

A number of investigators have surveyed psychiatric disorder present in patients in non-psychiatric beds. Lipowski² estimated that psychiatric disorder or distress of significant degree was present in 30% of the patients he studied. Bergmann and Eastham³ published a series of 100 elderly patients admitted to an acute medical unit in the UK, whom they screened for psychiatric disorder. They found that 7% had a diagnosis of dementia, 16% delirium and 19% functional illness. Mezey and Kellett⁴ summarized a series of UK studies and found prevalence of 5–51%.

Nowhere are such numbers of patients referred for a psychiatric opinion. Other studies have investigated consultation rates.

CONSULTATION RATES

Wallen⁵ and her co-workers looked at consultation rates in short-term general hospitals in the USA, using a national sample of 327 hospitals. This was retrospective and adequate information was not recorded in 25% of the hospitals. Patients admitted specifically for a psychiatric illness were excluded. Less than 1% of those admitted were referred for psychiatric opinion. The highest rates were in hospitals attached to medical schools, those in urban areas and those in the north-eastern USA. Not surprisingly, these characteristics were highly correlated with each other.

Wallen and co-workers found that, in general, female patients and younger patients were more likely to be referred. Those referred were sicker, i.e. had been given more "medical" diagnoses, and had more complex problems. They therefore

tended to use more resources. Ethnic origin was not a significant variable, but payment system was. Patients on Medicaid (government financed medical care for low-income persons) were more likely to be referred than those admitted under private insurance schemes.

In this, as in many earlier studies, all ages were considered together; in subsequent work, consultation rates for elderly patients were compared with those for younger patients. For instance, Popkin *et al.*⁶ compared a series of 266 psychiatric consultations to patients aged 60+ with consultations to a younger group. They found that the consultation rate for patients under 60 was 2.85% and for those aged 60+ 1.99%, a highly significant difference. In no specialty service did the rate for those aged 60+ exceed that for younger patients. The diagnoses given by psychiatrists differed between older and younger patient groups: 46% of the older group were diagnosed as having organic mental disorder compared with 14% of the younger inpatients. The psychiatrists also recommended more psychotropic medication for the older patients, which could be explained by the high percentage of organic diagnoses, and more diagnostic tests, which could not. The psychiatrists assessing the elderly patients were not described as specialists in geriatric psychiatry.

In the UK, consultation rates were also studied. At Guy's Hospital, Anstee⁷ looked at the pattern of referrals in 1968–1969—10 years after Fleminger and Mallett⁸ had done so. He found that the referral rate had doubled from 0.7% to 1.4%. Of the 254 patients referred, 49 were elderly; 35 had been referred from medical and 13 from surgical wards. Also at Guy's Hospital, Poynton⁹ compared referrals from August 1982 to November 1983, when there was no specialist psychogeriatric consultation service, with those from December 1983 to January 1985, when such a service had been introduced. The rate of referral rose from 0.64% to 1.40%, there being a greater rise in male referrals (0.34% to 0.96%) than in females (0.97% to 1.17%). In both periods, depression was the commonest single reason for referral. Anderson¹⁰ reported a similar rise in referral rate following the setting up of a specialist consultation service in Liverpool. The rate rose from 0.7% to 1.96%; proportionately more referrals were for depression in the second period (p. 142).

WHY SUCH LOW REFERRAL RATES?

Goldberg¹¹ suggests that medical and surgical patients are referred for psychiatric opinion either because some cue alerts their doctor

that there is a psychiatric problem or because the symptoms cannot be accounted for by known organic disease. He has suggested five reasons for non-detection:

1. Many patients do not provide cues, although they will describe their problem if asked.
2. Patients often mention depression or anxiety at the beginning of the interview, together with somatic symptoms, but only the latter are selected for further questioning.
3. Medical histories are often taken in open wards, where lack of privacy does not encourage patients to mention psychiatric problems.
4. A known organic cause does not exclude a psychiatric disorder.
5. Even when a psychiatric disorder is suspected, many clinicians are not confident of their ability to make a psychiatric assessment.

Other reasons for non-detection/non-referral that have been postulated include the following:

1. Attitudes of physician and of patient/patients' relatives. Physicians may still cherish misconceptions that depression and cognitive impairment are an inevitable accompaniment of ageing or that psychiatric disorder is not amenable to treatment. Fauman¹² found that 19% of surgeons and 9% of physicians believed that psychiatric illness was incurable. Elderly patients and their relatives may have a greater fear of public exposure and stigma than younger ones, although this may be changing.
2. Alternatively, some physicians may have an increased tolerance of cognitive and behavioural disturbance in their elderly patients.
3. Cognitive impairment, or symptoms such as paranoia, may make it hard for patients to relate their symptoms.
4. Some doctors feel that organic mental disorder is not a psychiatric problem.
5. Psychogeriatric services may be inaccessible, either geographically or personally, i.e. if too much pressure on the psychogeriatrician delays liaison visits.
6. There may be fear that a psychiatric referral will lead to an increased length of stay, either for further investigation or for treatment.
7. Physicians may wish to treat their patients' psychiatric problems themselves. They appear particularly likely to wish to treat depression, anxiety, psychosomatic disorders and organic brain disease, and most unlikely to wish to treat suicide attempts or psychoses¹².

The commonest reasons for referral are generally for advice on management, including drugs, or assessment of cognitive state. Other reasons include delayed recovery, advice on the psychiatric side effects of drugs, help in the selection of patients for other procedures, "disposal" problems or because of a previous psychiatric history. Covert reasons may emerge, e.g. a history of conflict, either between members of the medical team or between team and patient, especially when the patient declines to follow medical advice. The psychiatrist may then be expected to act as mediator.

There is no doubt, also, that referral rates may depend on local or personal factors. In one hospital, a high referral rate was thought to be due to the siting of the psychiatrist's office next to the medical ward, resulting in many verbal referrals.

Benbow¹³, reviewing the old age psychiatry service in the centre of a large conurbation, found that most referrals from hospital

doctors (58%) came from physicians in geriatric medicine, the remainder from other physicians and surgeons. These referrals accounted for 35% of the department's work load.

OUTCOME

What is the effect of referral to the psychiatry of old age service? In terms of immediate outcomes, the most frequent consequences appear to be suggestions for further investigation, advice concerning psychotropic medication, social services, counselling or other brief psychotherapeutic intervention or behavioural approaches by ward staff. Few patients are judged to need transfer to a psychiatric bed—6.6%¹⁴, 9%¹⁵.

Querido¹⁶ found that there was a significant increase in the chance of recovery for patients looked after by a team consisting of a physician, a social worker and a psychiatrist, assessing all aspects of their care, compared with those treated in a conventional way. He found that predictions based on team assessments were more accurate than clinical forecasts based on the patients' physical illness.

Others have shown that psychiatric disorder present during an inpatient stay is associated with subsequent increased mortality rates; e.g. in open heart surgery¹⁷ or from myocardial infarction^{18,19}. Treatment might be expected to have a positive effect on the mortality rate. It has also been shown that psychiatric intervention can lessen length of stay^{20,21}.

Hawton²², following up a cohort, examined during admission by Maguire *et al.*²³ after discharge, found that the presence of a psychiatric disorder during the initial admission was associated with higher subsequent mortality. He found that increased mortality rate was associated with age (not surprisingly) and the presence of psychiatric disorder. Excess mortality was not just due to more deaths in hospital or shortly after discharge, neither could it be explained by a higher percentage of patients with organic mental disorder; the mortality rate was also higher in patients with affective disorder.

Cooper²⁴ screened 626 patients aged 65–80 in an urban West German hospital and followed them up 1 year later. He found that the outcome—whether measured in terms of mortality, dependency or admission to continuing institutional care—was worse for those with organic mental disorder, 42% of whom had died at follow-up; 18% of those with functional illness, and the same percentage of those with "normal" mental states, had died. Functional illness appeared to correlate with increased dependency on others (42% needed help, as against 29% of "normals").

Feldman *et al.*²⁵ obtained similar findings: 49% of patients aged 70+ with organic mental disorder died within a year of assessment, compared with 20% of those without cognitive impairment.

Johnston *et al.*²⁶, in a prevalence study of psychiatric disorder in patients aged 65+ in non-psychiatric beds in a district general hospital, found that patients with significant psychiatric disorder had a greater length of stay. This appeared to be largely due to problems in placement.

Although some work has been done on the effects of intervention, less attention has been paid to financial aspects, either due to psychiatric intervention, or saved by it. This is becoming of vital importance to everyone in health care. Levitan and Kornfeld²⁷ studied outcome in a group of 24 patients undergoing surgery for fractured femur, where there was extensive liaison, and compared it with outcome in 26 patients treated in the same surroundings but without involvement of a psychogeriatrician. Mean length of stay for the first group was 30 days and for the control group 42 days, a significant difference. Twice as many

liaison patients returned home, as opposed to discharge to institutional care.

A SPECIALIST PSYCHOGERIATRIC LIAISON SERVICE?

Evidence has been cited to show that there is much undetected psychiatric disorder in patients occupying non-psychiatric beds, that there is a higher mortality rate in patients with such disorder, and that psychiatric intervention can reduce length of stay. Even though it would be wrong to assume, as some psychiatrists do, that all patients with emotional disorder require specialist assessment, it is probable that a good deal of suffering on the part of patients and their relatives and carers could be avoided by early detection and treatment. There is therefore a case to be made for a liaison service to elderly, as well as to younger, patients: "Coordination of medical care with any psychosocial care is likely to result in a better outcome and more effective use of medical services"²⁵.

Two models of service have been described: a "consultation" service and a "liaison" service. In a "consultation" service, the psychiatrist will only see patients specifically referred. Because of pressure of work, many of these assessments take place outside normal working hours and therefore without direct contact with the referrer. As a rule, further investigation and management will be recommended and will be carried out by the referring team. There is frequently no further contact with the psychiatrist unless problems occur—"please get in touch with me again if necessary...".

In a liaison service, a member of the psychiatric team joins the medical or surgical team usually on a weekly basis, e.g. on a ward round. He/she will take an active role in the further management of the patient. In practice, geriatric medical services most frequently have an attached liaison psychiatrist and many services offer a mixed consultation/liaison model.

Swanwick *et al.*²⁸ compared the two models. They found no major differences in reason for referral, broad diagnostic category or in suggested intervention and follow-up. They did find that in the liaison model there was a higher percentage of accurate diagnoses made by referring physicians.

Concern has often been expressed that the introduction of such a service would lead to too great an increase in referrals for the capacity of the old age psychiatry unit. Swanwick *et al.*²⁸ found no evidence for an increased referral rate—in fact, in the liaison model there were fewer referrals relative to the number of beds.

In other services, an increased referral rate has been described. Scott *et al.*²⁹, after setting up a liaison service with a senior psychiatric trainee (a specialist registrar) attending geriatric medicine ward rounds, found a 100% increase in the referral rate initially. Referrals from medical teams increased, while those from surgical teams remained the same, at a small percentage. This, however, was counteracted by the benefits achieved. Like Swanwick, she found an increased accuracy in diagnosis, 28% of patients referred having a diagnosis of affective disorder as against 12% before the service was introduced.

Studies have generally found that, with an increased knowledge of psychiatric disorders and the potential beneficial effects of treatment, there are fewer inappropriate referrals and physicians treat more such disorders themselves. Scott²⁹ and Baheerathan and Shah³⁰, who also studied the introduction of a liaison service, found that the benefits of change in referral pattern more than offset the cost of the liaison psychiatrist.

Most of the consultation/liaison work described has been carried out by psychiatrists. Collinson and Benbo³¹ describe a successful liaison service using an experienced psychiatric nurse. Others, for example Camus *et al.*³², advocate the benefits of a full

multiprofessional psychiatric team working within the general hospital with patients in non-psychiatric beds.

Why should this not be part of an overall liaison service, which might have organizational advantages? What can specialist psychogeriatricians offer? This has been answered by Popkin *et al.*⁶; they will have special knowledge and awareness with respect to the evaluation of physical illnesses that give rise to psychiatric symptoms; they are practised in differentiating organic mental disorder from mood disorders; they have extensive knowledge of psychotropic medication and of the consequences of altered pharmacokinetics in elderly people; and they are well equipped to evaluate and address psychosocial factors, the family and social support. Some of the aims of a psychogeriatric liaison service are listed in Table 134.1.

What will be the problems of providing such a service? They are likely to include:

- The identification of need.
- Accessibility of interested psychogeriatricians.
- Acceptability to patients, relatives, physicians/surgeons and members of their teams.
- Fear of increased length of stay for diagnosis and/or treatment.
- The cost—and who should pay.

With respect to identification of need, much can be done in educational programmes with all (not only medical) staff. In fact, most psychiatric disorder will be suggested by an adequate medical and social history and examination, as is taught in undergraduate curricula. Often the importance of this is not appreciated, and time and other factors do not encourage it. Other staff—nurses, occupational therapists, physiotherapists and social workers—are well able to identify and describe psychiatric problems if they are taught what to look for. They are also able to initiate plans for discharge at an early stage, e.g. with orthopaedic patients, rather than waiting until surgical rehabilitation is complete. Psychiatric staff need to learn the needs of their colleagues on the general wards and how to assist them; in particular, the fear and distress that mental illness and behavioural disturbance arouse in those unaccustomed to them. It is thus imperative that psychogeriatricians be involved in planning in-service training.

The use of screening tests has been shown to increase detection rates. The subject has been discussed extensively in recent papers^{26,33,34}. Geriatricians have found the abbreviated Mental Test Score³⁵ useful for cognitive impairment. The Mini-Mental State Examination³⁶ and the Clifton Assessment Procedures for the Elderly³⁷ have some advantages, the latter being rather long but more inclusive, the former quicker and therefore more acceptable. It is more difficult to single out tests for functional illness: brief versions of the General Health Questionnaire^{38,39} or the Geriatric Depression Scale⁴⁰ have been shown to be useful.

Table 134.1 Aims of a psychogeriatric liaison service

-
- To increase awareness of the prevalence and benefits of treating psychiatric disorders in elderly people
 - To stress availability of psychiatric advice
 - To provide a well-published referral system
 - To offer a prompt response
 - To offer a clear, practical assessment and recommendations with explanation, and transfer if needed
 - To adopt a helpful attitude to behavioural disturbance
 - To provide follow-up as inpatient and after discharge, if necessary
 - To run an acceptable educational programme
 - To undertake an evaluation of service
-

PSYCHIATRY OF OLD AGE UNIT: REQUEST FOR OPINION	
Patient name:	
Address:	
General Practitioner:	
Consultant:	
Urgent: Yes/No	
Date of birth:	
Telephone No:	
Hospital/Ward:	
Known to: Social Worker? CPN? Other?	
Reason for admission:	
Reason for referral:	
Relevant pathological findings:	
Screening tests: Mobility? Vision? Hearing?	
Living circumstances: Alone?	
	With carer?
	Other?
Social problems?	
Previous psychiatric history?	
Any other relevant information?	
Name of referrer:	
Position:	
Date:	

Figure 134.1 Specimen referral form. CPN, community psychiatric nurse

The Brief Assessment Schedule⁴¹ has been used to screen for both organic and functional disorder.

The majority of psychogeriatricians in the UK are aware of the importance of liaison; their accessibility varies. The organization of the service depends on local factors; many liaison services are based on linking a psychiatric team with one or more specialist firms. This has worked for much of adult psychiatry and has benefited both teams. It may conflict with the more usual pattern in psychiatry of old age services, where the team is responsible for residents of a defined catchment area and “follows” them throughout their care.

Those in other specialties need to know of the psychogeriatric liaison service. Notices in ward information packs and inclusion in induction courses for new staff are essential. Often the provision of a special referral sheet, such as that in Figure 134.1, helps both to remind and to focus on the information needed by the psychiatrist. There must be a means of telephone referral—preferably one number, with the psychiatric department sorting out the details of who should deal with the referral. As there must be an emergency service available out of hours, and this is usually provided by the duty psychiatrist, he, too, needs instruction.

Acceptability is usually best achieved by the results of an effective service. For instance, Bergmann and Eastham³ stated that, with respect to affective disorder, when liaison work took place staff attitudes on medical wards changed, including “. . . a new enthusiasm and willingness to look for affective disorders in other patients and to try to obtain treatment for them whenever possible”.

Similarly, fear concerning length of stay and cost can best be assuaged by explanation and discussion, and finally resolved in the light of successful practical experience.

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Education and the Liaison Psychogeriatrician

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The elderly are under-represented among referrals to psychiatry from general hospital wards in relation to younger patients and bed occupancy. Less than 3% of general elderly admissions are seen by a psychiatrist and only 20–25% of all liaison contacts are with older people, even though they occupy 40–50% of general hospital beds and 30–50% might be expected to have or develop a psychiatric problem¹. This represents 3000–5000 cases/10 000 admissions and even if more were referred, only a fraction could realistically be seen by a psychogeriatrician².

The frequency of mental disorder in acute elderly medical admission populations is approximately twice that of the community^{3,4}. Depression, dementia and delirium are the common syndromes. Psychiatric problems are much more common on medical and orthopaedic than general surgical wards^{5–7}. The frequency of organic brain syndromes increases with age⁸ but there is less evidence that depression is age related⁹.

The identification of mental disorder by general clinicians is poor¹⁰, particularly the detection of depression¹¹ and delirium^{12,13}. Clinicians appear to detect the syndrome of dementia more reliably than cognitive symptoms¹⁴ but with depression it is symptoms that are more often recognized than the syndrome¹⁵. The possible adverse effects of a co-morbid psychiatric disorder upon the outcome of a medical admission makes the recognition and treatment of these disorders important to both patient and clinical service^{6,16,17}.

A priority for the psychogeriatric consultation–liaison service has to be education (acting as it does at the interface between psychiatry and general departments) that encourages good practice and alters attitudes to mental illness in old age. With such morbidity it is essential to improve the ability of general staff to detect and prescribe appropriate treatment for the majority of simple disorders, while recognizing those cases that need the specialist psychiatric service¹⁸. Currently, the treatment of mental disorder by general staff, particularly for depression, appears poor¹⁹.

There is indirect evidence that specialist consultation–liaison psychiatry for the elderly does influence the behaviour of general clinical staff toward psychiatric problems. The introduction of a consultation–liaison service is certainly associated with increased rate of referral^{1,20–23}. This increase is most marked for depression^{1,23}, perhaps the most neglected and inappropriately managed condition, but close liaison produces a general improvement in the quality of referrals²³.

Research in the old age liaison field is in its infancy and has concentrated on quantifying levels of morbidity, examining referral rates and exposing clinicians' difficulties in recognizing psychiatric disorders. Preliminary work suggests that psychiatric involvement with older patients can have positive effects on

outcome²⁴ but this may not be targeted at the most appropriate cases¹⁸. If the management of mental disorder in this context is to improve, then general services will need education. The process of education is complex and multifaceted and a new research direction would involve a closer examination of the enabling role of specialist consultation–liaison, identifying approaches and style of service with the greatest educational impact and the most effective methods of disseminating knowledge and expertise.

It is the management of non-referred cases that will ultimately prove the measure of success. As the ageing population grows, this research is timely and these important areas of study need to be explored.

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Rehabilitation

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Rehabilitation of the older person with psychiatric disorder means restoring and maintaining the highest possible level of psychological, physical and social function despite the disabling effects of illness. More broadly, it also means preventing unnecessary handicap associated with illness, preventing unnecessary handicap secondary to maladaptive responses to illness, and combating the deadening effects of low expectations of older people amongst patients, families and society in general. Managing chronic disease and disability is the greatest challenge to modern medicine. Within this, the rehabilitation of many older people with psychiatric disorder looms large—although, of course, many old people with psychiatric disorders respond well to “curative” therapy and require little rehabilitation.

In fact, rehabilitation is a fundamental and inseparable part of old age psychiatry. Perhaps for this reason, as with rehabilitation in geriatric medicine¹, little has been written about the topic specifically. Some particular techniques, such as psychological approaches with the cognitively impaired, have been well described^{2,3} but little evaluated⁴, and evaluative research is much needed here.

SPECIAL PROBLEMS WITH PSYCHIATRIC DISORDER IN THE ELDERLY

“Old age” may span 30 years or more, posing quite different rehabilitation problems; but the most major concern is with the old-old. In this group, multiple disability is prominent, with the complicating danger of polypharmacy, and physical and mental ill-health interact in complex ways. With this frail population, disentangling the respective influences of ageing, previous personality and current ill-health can be exacting. Two-thirds of the UK's disabled population are older people and the true extent of handicap due to psychiatric disorder is probably still not established.

With depression, especially, there may be restriction of physical activity, threatening physical capacity and health. Depression associated with stroke disorder⁵ or with Parkinson's disease⁶ particularly illustrates both the connection between physical and psychiatric problems and the importance of physiotherapy in psychiatric rehabilitation.

Physical factors are frequently of great importance in dementia. A quiescent individual may become delirious and disturbed at night through heart failure, obstructive airways disease or even the uncomfortable effects of severe constipation. Settling such problems may transform the reality of care for a carer and seeking out such therapeutic opportunities is an important part of rehabilitation. Similarly, in dementia, physiotherapy to promote and maintain the best possible physical capacity is a key element.

Advice and practical aid to carers, such as with lifting and handling the physically disabled demented person, can be crucial.

A judicious mixture of the “therapeutic” (curative) and the “prosthetic” (supportive) approaches^{7,8} is very necessary in old age psychiatry. Whilst much functional psychiatric disorder and delirium can be “cured”, and this must be the aim, most older people with dementia need some degree of supportive care at some stage. The poor financial and housing state of many older people, together with the lack of children or spouses to help as carers for many of the old-old, are further complicating factors. Maximizing “participation” despite psychiatric disorder needs to be a major goal, maintaining as far as possible a role in the family, social contact, a range of activities and a minimization of loss of autonomy or institutionalization. This requires an approach that embraces psychiatric, medical, rehabilitation, nursing and social perspectives. Seeking active prevention of disability/reduced participation as a consequence of psychiatric disorder is a vital part of rehabilitation.

SPECIAL PRINCIPLES IN REHABILITATION OF THE OLDER PERSON WITH PSYCHIATRIC DISORDER

Table 136.1 summarizes the principles. The first principle is to make the home the focus of attention. That is where problems have arisen and where they will need to be overcome. Planning for rehabilitation should begin at the earliest moment; an initial assessment at home by a senior psychiatrist, or other experienced team member, is invaluable—even if “home” is an institution in the community. Home is also where the carers, and often any social services support staff involved, may readily be found. This contrasts with standard medical rehabilitation, where often no such opportunity exists.

Table 136.1 Principles of rehabilitation for the older person with psychiatric disorder

Focus on the home
Ensure comprehensive assessment
Encourage normal function
Treat the treatable
Analyse disabilities and chart progress
Clarify team goal with patient and carers early
Clarify team goal with support workers early
Teach what can be learnt
Adapt the adaptable
Coordinate support and follow-up
Promote flexibility and ingenuity
Promote realistic optimism

Frequently further assessment by, say, an occupational therapist, physiotherapist or another specialist team member may be necessary—and this can perhaps also be arranged at home. But avoiding disruptive admission should not be at the cost of meeting assessment needs. The day hospital and its team is often useful to complete such assessments.

Poor function or morbidity should never be accepted as immutable and still less as normal. Assessment should aim at a thorough understanding of social, physical and psychological function, as well as the previous pattern of personality and lifestyle. From this, diagnoses and specific treatment for the treatable should follow. But also analysis should show the extent of disability, how it is mediated and how it may be overcome. Problems should be clearly recorded, with proposed solutions and with regular review of progress^{9,10}. From the earliest moment, independent function and improvement should be sensitively encouraged, especially with hospitalized patients.

As early as possible, the agreed goal of rehabilitative efforts needs to be clarified with the patient, with the relatives/carers, with the whole rehabilitation team and with any support workers needed in the community. Education, guidance and a rehabilitative “demonstration” may well be necessary to resolve conflicting views on the prospects for progress. Carers may need therapy or rehabilitation in their own right. Almost always the support of carers is essential, although some stoutly independent patients manage well without this.

A prognosis-based plan is useful. This means assessing: what are the problems and what are their causes? Prognostication will follow: will these get better or worse? To what degree? Over how long a period? From this a plan can flow consistent with what seem the most major likely developments of a problematic nature, over a foreseeable timescale and taking account of what is most remediable.

Sometimes, disability strongly distorts the previously stable power and dominance pattern in the family¹¹ so that, for instance, a forceful mother becomes dependent on a passive daughter. Relationships are always important and such phenomena can strongly influence outcome. They need to be understood and often complex ambivalence worked through¹². Such factors are frequently important when carers seem reluctant to resume caring¹³ and need to be carefully teased out when juggling with the various elements of risk and risk minimization in supporting a vulnerable person at home¹⁴. The patient’s “crutch is not made of wood but of some other person’s tolerance or patience”¹⁵.

An agreed balance should be sought between the needs of carers and the patient’s right and desire for continued comparative independence despite significant disability—bearing in mind the

team’s prime responsibility to the patient. In effect, carers should be “recruited” as rehabilitation therapists’ “aides”. But often they will need practical help and advice on rehabilitation techniques and, always, the reassurance of the services’ continuing availability for support and sensitive expert response. Similar considerations apply with any support staff necessary to help the patient at home—principally social services staff in the UK. Their early involvement and integration into the assessment and rehabilitation process can be logistically difficult but generally is most effective. Good teamwork is of the essence. Clarity and consistency within the multidisciplinary specialist team are essential. Nurses and therapists, for instance, must communicate well, each complementing and enhancing the other’s approach. Specialist and general practitioner must be in accord. The specialist team must carry the confidence of those who will work with the patient outside hospital. Table 136.2 lists many of the team members and resources requiring coordination for rehabilitation—inevitably there is great overlap⁷. Clear goals should be set, in accord with patient and carers, and clearly understood by all^{16,17}. This is much easier said than done with the vulnerable frail elderly person; but vitally important is good communication.

Patients are taught what they can learn or relearn but often only modification of domestic equipment, provision of aids or modifications of the home will overcome their disability. More often still, problems are only overcome through support services—home-delivered meals or a home help/community care assistant—coming into the home. Ingenuity and diplomacy may be needed with an old person reluctant to accept such necessary support. The great majority of support is provided by relatives/carers and supporting the supporters is the main task. This may require day hospital therapy, day care or respite admissions. Maintaining confidence in the care service is vital.

At some stage with hospital patients, except in the most grossly deteriorated person, a home assessment is advisable with an occupational therapist or physiotherapist, or both, and perhaps with other team members. Hospital-based staff can be too pessimistic and, allowed to function in his/her familiar environment, even a significantly demented patient can sometimes perform surprisingly well. Often serial home assessments are helpful with increasing challenge, leading to overnight stays. Also, initial failure to manage satisfactorily should not preclude the possibility of later improvement with further therapy or support.

Above all, rehabilitation with older people requires flexibility allied to a realistic but constructively optimistic approach. Innovation and ingenuity are frequently necessary. The complex interaction between physical, social and psychological factors in older patients is further complicated by the likelihood that

Table 136.2 People and facilities to aid rehabilitation of the older person with psychiatric disorder

	Specialist psychiatric team	Social services and local authority services	Primary care team
People	Psychiatrists Ready access to geriatricians Hospital nurses Community psychiatric nurses Physiotherapist Occupational therapist (OT) Clinical psychologist Social worker (Speech therapist—sometimes) (Dietician—sometimes) Carers’ groups	Social workers Domiciliary services manager Home help/community care assistants Meals on Wheels Community OT Support to voluntary sector Sitter services Carers’ groups	General practitioners Practice nurse District nurse Health visitor
Facilities	Assessment ward Nursing home Day hospital Outpatients Long-stay and respite care facilities	Luncheon club (often voluntary) Day centres (often voluntary) Long-stay and short-stay residential care Housing adaptation Sheltered housing	Health centre

circumstances, perhaps especially physical health, may change dramatically; this demands a flexible response. The solution, carefully constructed and successful on one day, may need major change on the next as the situation radically alters. For this reason, rehabilitation with older people is rarely completely finished and "maintenance" measures are often necessary. Careful planning of continuing care, follow-up and continuing availability as problems arise are essential features. The strength and determination of patients, relatives and support staff are considerably bolstered by the knowledge that this approach is backing them up.

REHABILITATION AND LONG-STAY CARE

Many patients, predominantly demented individuals, will require long-term care but rehabilitation must remain a strong theme. Such deteriorating multiply-disabled demented patients require 24 h care. They often exhibit difficult and disturbed behaviour and need heavy physical care. Providing the best quality of life, given often quite limited resources, is the aim and a major strategy is preventing unnecessary dependency and promoting the maximum retention of function. With descriptions¹⁸ of the care of such patients in various settings, a frequent theme has been the availability of skilled and expert staff from the multidisciplinary team to help maintain good function. Evaluation of such long-term care programmes has proved complex and difficult^{19,20}.

There has been concern in the UK that such disabled patients will increasingly be excluded from hospital care for funding reasons, in favour of health authority contracted private nursing home care or simply standard care in private nursing homes²⁰⁻²². In the USA (and many other countries), much long-term care of older people has long been provided in the private sector. But in the USA the Omnibus Budget Reconciliation Act (OBRA Act)²³ requires nursing homes to ascertain and meet any needs for therapy and treatment. In the UK the fear is that the drive in the best hospital care to provide a good quality of life may be replaced in private nursing homes by a desire for a quiet life; passivity and dependency (possibly resulting from unnecessary tranquilizing medication²⁴⁻²⁶) could be more acceptable than the patient's exercise of individuality, movement and self-expression.

Reports of high levels of depression in homes for older people²⁷⁻³⁰ emphasize the worry about effective rehabilitation and care for older people with chronic psychiatric disorder. Any institution providing shelter is at risk of providing a relatively impoverished environment¹⁵ and undue restriction of independence.

Bennett¹⁵ called unnecessary social inactivity and dependence "the psychiatric equivalent of contractures". Hospitals (and social services) have provided much good practice. Accounts of good long-stay care for demented people emphasize the invaluable input of occupational therapy, physiotherapy and person-centred approaches^{30,31}. Sensibly ensuring activity in a structured day and enriching the environment with, for example, music therapy, art therapy, drama therapy³² or reminiscence therapy³³ are important wherever the setting.

Community psychiatric nurse (CPN) support (indeed, availability of all the specialist team to give support) to such homes can be feasible and helpful. But concern remains about how to monitor effectively and maintain standards in private long-stay care²⁰. Ultimately, good care here depends on the commitment of sufficient appropriately trained staff to help disabled old people experience to the full their remaining scope for independence and capacity for joy.

ATTITUDES

Adverse feelings about older people have been noted by many³⁴. Geriatricians have commented on unhelpful attitudes in health

professionals of all kinds, including GPs¹³. Modern medical education is inculcating a better knowledge base for tomorrow's doctors and also better attitudes³⁵. The educational potential of psychogeriatric services, not least exploiting the educational opportunity afforded by rehabilitation (and other concerns) with the most disabled, has been described³⁶. Not only medical students³⁷, but all varieties of medical and other professional staff concerned with older people with mental health problems, and also carers and lay audiences, benefit from this educational effort. Public education especially is vital in engendering constructive attitudes in society, on which ultimately will depend all efforts towards care in the community and the political will to provide decent services.

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Anaesthetics and Mental State

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The anaesthetist uses many drugs that directly or indirectly affect mental status and, as is discussed below, age-related changes in drug handling tend to increase the risk of adverse cognitive effects. The multiplicity of drugs used in modern anaesthesia also increase the risk of drug interactions, and at the same time make it more difficult to decide the cause of any adverse effect that occurs. For instance, a typical general anaesthetic might involve premedication drugs, intravenous induction agents, gases and volatile agents, competitive neuromuscular blockers, drugs to reverse the neuromuscular blockade, opiate and non-opiate analgesics (which may be given pre-, intra- or postoperatively) and major and minor tranquillizers.

It might be thought that a "regional" anaesthetic (such as a local, spinal or epidural anaesthetic), where the patient maintains consciousness, would be free from central nervous system effects. However, premedication is often given prior to a regional anaesthetic and sedatives and analgesics may be administered during and following the operation. All of these may have a prolonged effect on mental status. Regional anaesthesia might also influence postoperative outcome, for good or ill, by affecting the ability to mount a "stress" response.

Apart from the expected pharmacological effects of anaesthetic agents on mental status, there are also unwanted reactions to be considered, such as the increased tendency of anticholinergic drugs to precipitate acute delirium in older patients. Secondary drug effects may also be important, such as the disturbance of sleep pattern by opiates, which has been hypothesized as a cause of late postoperative hypoxaemia. Many of these individual effects are discussed below.

It should also be appreciated that an acute or chronic change in cognitive state which follows an anaesthetic may have nothing to do with that anaesthetic. Acute delirium can be precipitated by a variety of "stresses", including acute illness and non-anaesthetic drugs, and also by withdrawal from drugs, such as alcohol and tranquillizers, which were being taken prior to surgery^{1,2}. Neurological damage following operation may result from hypotension or cardiac arrhythmias, which had nothing to do with the anaesthetic, hypocarbia may reduce cerebral blood flow, and the patient who appears demented for the first time following emergency surgery may have been dementing quietly and unobserved for many months before.

In an attempt to bring some structure to the above complexity, the following discussion will be divided into three parts: first, age-related changes in drug handling which are of particular importance to the anaesthetist; second, major direct and indirect mechanisms by which anaesthetic agents might affect mental function; and third, short-term and long-term studies of psychometric testing before and after anaesthesia (including the effects of clinical risk factors and type of anaesthetic, and the findings of the

ongoing ISPOCD studies). It should be recognized, however, that the list is not exhaustive. For example, an operation under anaesthesia, like any other major life event, may precipitate anxiety and other changes in mental status³.

Cognitive changes that occur after open heart surgery have also been well studied in the literature. However, in this group of patients, the cerebral effects of the cardiac bypass procedure add another layer of complexity to the analysis of the postoperative outcome, and they have not been further discussed in the present brief account, although part of the recent comprehensive review into postoperative cognitive dysfunction by Dodds and Allison⁴ deals with this subject.

AGE CHANGES IN DRUG HANDLING THAT ARE OF PARTICULAR RELEVANCE TO THE ANAESTHETIST

As a broad generalization, Dodson⁵ has subdivided anaesthetic drugs into two classes:

1. Drugs to which the elderly patient tends to demonstrate *increased sensitivity*. This class embraces most of the drugs that affect the central nervous system, including sedatives and analgesics. Such drugs usually have a more profound and long-lasting effect in the elderly. There may also be an increased incidence of side-effects of drugs in old age, such as muscle rigidity following fentanyl, and delirium following anticholinergic agents.
2. Drugs that act on the autonomic nervous system, where the elderly patient tends to demonstrate *decreased sensitivity* at receptor sites. Thus, the effects of autonomic blockade by anticholinergic drugs or β -blockers may be less dramatic, as may be the effects of vagal and sympathetic stimulants.

While the above classification is a useful conceptual framework, dosage regimens in individual elderly patients need to take into account individual differences in drug susceptibility, age-associated illness and physiological status. In most cases, however, the anaesthetist will need to give smaller doses of drugs to elderly patients^{5,6}.

In a more recent review, Dodds⁷ has described anaesthesia as "applied clinical pharmacology with enough patho-physiology to confuse the picture". He lists nine key points, three of which are of direct relevance to mental function, and six of which are of potential relevance:

1. The minimum alveolar concentration (MAC) is that concentration of an anaesthetic gas which suppresses movement in

- response to a surgical stimulus in 50% of subjects. MACs tend to fall with age.
2. Because of reduced neuronal density and a reduced metabolic rate, the elderly may be more sensitive to a given amount of anaesthetic drug. In elderly patients with a delayed circulation time, there is an added potential for overdosage when intravenous agents are given too fast, because a delayed response may be mistaken for a lack of therapeutic effect, so that when the drug eventually reaches the brain, too much drug has been given. Smaller doses, slower rates of infusion or repeated small boluses are recommended in the elderly.
 3. The elderly appear to have an increased sensitivity to opiates.
 4. Where there is pre-existing ischaemic heart disease, the patient may be vulnerable to the cardiac depressant effects of some anaesthetics. There is also a theoretical risk that some anaesthetic agents can "steal" blood from ischaemic myocardium by increasing vasodilation in normal vessels.
 5. With increasing age there is a tendency to increased shunting in the lungs and decreased cardiac output. These pathophysiological changes have complex effects on the uptake of volatile gases, as a low cardiac output favours rapid uptake, while a decreased lung function produces the opposite effect. Such effects are less with the more insoluble gases.
 6. Hepatic metabolism and/or clearance of drugs tends to be affected by age, although there is considerable interindividual variability. Insoluble anaesthetic agents which require no hepatic metabolism should be safer than soluble agents. Some anaesthetics, such as halothane, are also potentially hepatotoxic.
 7. Renal clearance of drugs tends to fall with age in some, but not all older people. Some anaesthetics, such as sevoflurane, are potentially nephrotoxic.
 8. Neuromuscular blocking agents can be classified into depolarizing types and non-depolarizing types. The non-depolarizing agents are usually favoured in old age.
 9. Non-steroidal anti-inflammatory agents are increasingly being used for analgesia in younger patients, but may present increased hazards in the elderly because of nephrotoxicity, fluid retention and tendency to gastric irritation.

MECHANISMS BY WHICH INDIVIDUAL ANAESTHETICS MIGHT AFFECT MENTAL FUNCTION

A fuller discussion of this topic can be found in the review of Dodds and Allison⁴, but major potential mechanisms are as follows:

Direct Cerebral Effects

Trace amounts of general anaesthetic agents can have effects on alertness and concentration after patients recover consciousness. Experiments involving young volunteers have also documented the effects of subanaesthetic doses of drugs on mental function, behaviour and motor skills^{8,9}, and have also studied the effects of prolonged general anaesthesia in the absence of surgery¹⁰. None of these studies involved old subjects, but it would be expected that the central effects of anaesthetics would become more marked with age.

Hypoxaemia

In recent years, pulse oximeters have provided a convenient, non-invasive means of monitoring postoperative hypoxaemia¹¹. The

new technology produced some unpleasant surprises: many elderly patients who appear to be stable and well-perfused experience profound and prolonged episodes of nocturnal hypoxaemia several days after surgery. Such episodes might explain some of the "unexpected" episodes of confusion, cardiac abnormality and sudden death that can occur up to a week following operation. However, as discussed in the review of Dodds and Allison⁴, some researchers are not convinced that hypoxaemia is a major cause of temporary or permanent postoperative cerebral dysfunction (POCD), and the ISPOCD1 Study (see below) failed to find a correlation between hypoxaemia and postoperative cognitive status, even though it set out with the hypothesis that such a relationship existed.

The mechanisms by which anaesthetic agents, particularly opiates, might cause postoperative hypoxaemia are reviewed by Jones and his colleagues¹¹. It appears that postoperative hypoxaemia is caused by the interaction of two major factors. The first of these factors is a gas exchange abnormality induced during anaesthesia, which tends to be most marked in the first few postoperative hours but which may persist longer. The second factor is episodic obstructive sleep apnoea, which may occur several nights after anaesthesia. The episodes of obstructive breathing seem to be precipitated by the *return* of rapid eye movement (REM) sleep, which had been abolished for a few days by drugs, particularly opiates, given around the time of operation, and by the stress of operation itself. This late hypoxaemia, once recognized, can be alleviated by giving oxygen, but in high risk patients a week of oxygen administration may be required.

Anticholinergics

As has already been explained, anticholinergic drugs tend to be less effective in blocking cholinergic function in old age. Much more important, however, are the direct central effects of anticholinergic drugs (such as atropine) that can cross the blood-brain barrier. In some cases excessive drowsiness results, while in others hallucinations and delirium occur, and some authors refer to a "central anticholinergic syndrome". A number of drugs that are not primarily anticholinergic agents also have anticholinergic properties, including phenothiazines, pethidine, thiobarbitone, flurazepam and tricyclic antidepressants. Two decades ago, in a small group of open heart surgery patients aged 29-75, Tune *et al.*¹⁰ reported that the presence and severity of postoperative delirium was correlated with the level of anticholinergic drug activity in the blood. More recently, O'Keeffe and Chonchubhair¹³ have reviewed the potential contribution of anticholinergic drugs to postoperative delirium, and have considered their importance relative to other pharmacological and non-pharmacological mechanisms.

While the evidence is not absolutely conclusive, it would seem prudent to minimize anticholinergic use in older surgical patients. One way to achieve this would be to avoid anticholinergics in premedication regimens, or perhaps to avoid premedication altogether. If an anticholinergic drug is thought essential, then glycopyrrolate, which does not cross the blood-brain barrier, would appear to have advantages over atropine¹⁴.

COGNITIVE TESTING IN YOUNG AND OLD PATIENTS BEFORE AND AFTER SURGERY

Postoperative Delirium

Before considering the more subtle cognitive effects that can occur postoperatively, the syndrome of acute postoperative delirium will be discussed. The causes of postoperative delirium,

as in delirium in other settings, include acute illness, drugs and drug withdrawal^{1,2,13,15} but the incidence tends to rise with age. Estimates of the incidence of postoperative delirium in the over-65s range between 7% and 50%, depending on the definitions used and the clinical circumstances^{2,13,15}. The incidence tends to rise with age, the urgency of surgery, the use of sedative and anticholinergic drugs and the degree of preoperative mental impairment. Factors (such as sepsis) that favour the development of delirium in a non-surgical situation¹² may also be of relevance postoperatively.

Recent advances in the study of postoperative delirium have included attempts to standardize definitions¹³ and a large study in the USA by Marcantonio *et al.*¹⁵, which attempted to develop a clinical prediction rule in 1341 patients aged 50+ having major non-cardiac surgery. The latter authors found that seven preoperative factors (age over 70, alcohol abuse, poor cognitive status, poor functional status, serum electrolyte/glucose disturbances, thoracic surgery and aneurysm surgery) had an independent relationship with postoperative delirium. However, the effect of intraoperative events, including anaesthesia, was not studied, as the aim was to produce a preoperative rule.

O'Keefe and Chonchubhair¹³ have concluded that, in at least 90% of cases of delirium following general surgery, it is postoperative medical or surgical complications that are to blame, which implies that the appearance of delirium should lead to a diligent search for underlying physical medical problems. The increased incidence of postoperative delirium with age is likely to be due to causes such as these, rather than effects arising from age-related differences in handling anaesthesia¹⁶.

Postoperative Changes in Psychometric Tests in Older People

Delirium is an important postoperative syndrome, but more worrying is the possibility that *dementia* could occur for the first time as the direct result of *routine* surgery and/or anaesthesia (as opposed to the mental changes that might arise from an intraoperative catastrophe, such as cardiac arrest). In 1955, Bedford¹⁷ published a much-quoted retrospective study in which he sought to trace those patients in the Oxfordshire area who had "never been the same again" after an elective or emergency anaesthetic. Over a 5 year period he was able to identify 18 cases where there was reasonable evidence that dementia had appeared for the first time after surgery and anaesthesia. In interpreting this data, it is important to realize that a detailed assessment of preoperative mental function was not available, and that even in cases where dementia was reported immediately following surgery, it did not necessarily imply that anaesthetic drugs were the cause.

In a subsequent study, Simpson *et al.*¹⁸ attempted the very difficult task of replicating Bedford's findings in a prospective study. As formal preoperative psychological assessment was part of Simpson's study design, only elective patients could be included. After considerable efforts, 678 elderly patients having surgery were evaluated, two-thirds of whom underwent general anaesthesia. This major undertaking yielded only one patient in whom there was good evidence that a permanent deterioration in mental function occurred immediately after anaesthesia.

These two reports^{17,18} stimulated many psychometric studies of short- and long-term postoperative cognitive outcome. Tables 137.1 and 137.2 summarize the results of 17 such studies¹⁹⁻³⁵, which have looked at psychometric test performance in ageing patients before and after surgery. Follow-up periods have ranged from a few days to over 3 months, and many different types of

psychometric tests have been used. Table 137.1 contains 13 studies in which older patients have been randomized to receive general or regional anaesthesia, while Table 137.2 summarizes four alternative study designs that have given an insight into the effect of age on postoperative cognitive problems. These include the ISPOCD1 study, which is discussed in detail below.

There are major methodological problems in carrying out clinically meaningful psychometric testing in elderly elective surgical patients³⁵. These problems become almost insuperable in patients who are admitted as emergencies, and only two of the studies listed in Tables 137.1 and 137.2 considered non-elective patients. Unfortunately, there is ample evidence that it is non-elective patients who have the highest incidence of postoperative mental events, although in many emergency cases non-anaesthetic factors, such as acute illness, are probably more culpable than anaesthetic drugs.

The 13 studies in Table 137.1 compared the psychometric effects of general anaesthesia with those associated with "regional" techniques (local, spinal or epidural anaesthesia). Four of the earlier studies reported that the general anaesthetic group performed more poorly, but even here the effect was seen only during the immediate postoperative period.

The studies that make up Tables 137.1 and 137.2 contain a wealth of detail, but some broad overall conclusions can be drawn. As might be expected, the major effects on mental function are seen in the first postoperative day, but some of the studies report minor effects on some tests for up to 7 days. The studies that specifically compared young and old patients found that the older group performed slightly worse than the younger during the early postoperative period.

While *short-term* mental impairment following surgery is of importance, especially in these days of increasing day surgery provision³⁶, it is *long-term* mental impairment that is feared most by patients, their relatives and their doctors. In the first edition of this textbook³⁷, comfort was drawn from the fact that the best-designed of the long-term studies up to that date had reported no objective evidence of mental impairment 1 month or more after surgery. It was noted, however, that 16% of the patients of Jones *et al.*²¹ had complained of subjective changes in memory and concentration at 3 months after surgery, and that these authors had commented that these patients might have had minor intellectual changes which had been missed by the chosen psychological tests. Since that time, the study of Williams-Russo *et al.*¹⁹ has similarly reported long-term postoperative cognitive changes in 5% of patients. Further concern has been raised by the finding of long-term cognitive deficit in the ISPOCD1 study³², which, unlike the Williams-Russo *et al.* study, included a group of non-operated control patients to meet the criticism that some of the effects reported in early studies reflected the progression of coincidental dementia, unrelated to surgery or anaesthesia. The ISPOCD1 study and its successor, ISPOCD2, will now be described in some detail.

The ISPOCD Studies

The first International Study of Post-operative Cognitive Dysfunction (ISPOCD1) collected data between 1994 and 1996 on 1218 patients aged 60+ who were undergoing major non-cardiac surgery in 13 hospitals in eight European countries and the USA³². This was a major undertaking, which was intended to answer many of the questions about early and late postoperative cognitive dysfunction in older people that had been raised in the literature over the previous 30 years. In the event, the results of the ISPOCD1 study, published in 1998³², still left several unanswered questions which are being addressed in a further study (ISPOCD2), which is due to report in May 2001.

Table 137.1 Randomized studies comparing the psychometric effects* of general anaesthesia (GA) and regional anaesthesia** (RA)

Reference	Number of patients	Age (years)	Type of surgery [anaesthesia]	Timing of postoperative tests	Main findings	Difference between GA/RA?	Long-term cognitive deficit?
19	262	>40, median 69	knee replacement (TKR) [Epidural vs. general anaesthesia]	7 days, 6 months	Delirium occurred in 11%. Complex changes in psychometric tests, but there was a general reduction at 7 days with return to baseline (or better) by 6 months. However, Trail Making A and B worse at 6 months	No	5% Showed significant deterioration at 6 months, but there were no untreated controls
20	169	65–98	Cataract [GA vs. local]	1, 14 days, 3 months	Reduction in verbal recall/learning, psychomotor speed and tactile naming on day 1 only	No	No
21	146 patients, 50 controls (patients on waiting list)	60+	hip (THR) or knee replacement [GA vs. epidural]	3 months	No decreases on tests in patients or controls at 3 months. Tests included Choice Reaction Time (which <i>increased</i> in GA group), object learning, digit copying, critical flicker fusion, and cognitive difficulties scale	No	No <i>measurable</i> decrease, but 11/56 GA and 10/60 RA patients thought concentration and memory were poorer
22	64	60–86	TKR [GA vs. spinal]	3 months	Tests tended to improve at 3 months (? practice effect)	No	No
23	40	60–80	Transurethral resection of prostate (TURP) [GA vs. epidural]	4, 21 days	Day 4 only: reduced paired associate learning, visual memory and visual recall	No	No
24	105	25–86	Hysterectomy, TURP, THR/TKR	1–7 days, 3 months	Modest decrease in memory and cognitive tasks in early period. Later tests usually better than baseline	No	No
25	30	50–80	THR [GA vs. spinal]	1, 2, 7 days	Spinal group had worse word recall/recognition on day 1, no differences on later days	Yes	Not tested
26	44	60–93	TURP, pelvic floor surgery [GA vs. spinal]	6 h, 1, 3, 5 days, 1 month	At 6 h GA group had reduced Mini-Mental Status. No later differences on Mini-Mental or Geriatric Mental Assessment	Yes	No
27	57	65–92	Hip fracture [GA vs. epidural]	1, 7 days	38–50% Delirium (even though patients excluded if Organic Brain Syndrome Scale abnormal preoperatively)	No	Not tested
28	40	>60	Hip fracture [GA vs. spinal]	7 days, 3 months	Abbreviated Mental Test <i>improved</i> at 1 week (no other test was used)	No	No
29	30	>60	THR [GA vs. epidural vs. both]	2, 4, 7 days, 3 months	Assessment by psychologist. Various tests impaired for 2–4 days, then recovered	No	No
30	60	>65	Cataract [GA vs. local + sedation]	7 days	Wechsler Memory Scale and Luria tests reduced relative to baseline	Yes (LA <i>worse</i> , Luria only)	Not tested
31	60	56–84	THR [GA vs. epidural]	1, 3, 7, 12 days	No formal psychological tests; 7 out of 31 in GA group (0 out of 29 epidural group) said to have mental changes in first 7 days	Yes	5 out of 31 GA patients reported mental changes 4–10 months after surgery

*For full details of psychometric tests, see references 4, 37.

**Regional anaesthesia includes local, spinal and epidural anaesthesia.

A key feature of the ISPOCD studies has been the use of the European psychometric test battery (EUPT battery), which has been designed as a sensitive and standardized research tool for the detection of postoperative neuropsychological deficits and which can be administered over a 45 min period. This battery uses the Mini-Mental State Examination (MMSE) as a screening test, with patients scoring 23 or less being excluded from further testing. The remaining tests (used in ISPOCD1 and in a slightly modified form in ISPOCD2) comprise a Verbal Learning Test, a Concept Shifting Test, the Stroop Colour Word interference test, a Letter

Digit substitution task, a four boxes test, the Broadbent Cognitive Failure Questionnaire, and a Zung Depression Score Questionnaire (the Geriatric Depression Scale being used in ISPOCD2). The definition of postoperative cognitive dysfunction (POCD) has been reached by comparing changes in the normalized (Z) scores of individual patients with age-matched controls who were not undergoing surgery, who were studied at the same time intervals.

ISPOCD1³² was particularly concerned to test the hypothesis that hypoxaemia and/or hypotension were causative factors in POCD. Accordingly, oxygen saturation was measured by

Table 137.2 Other studies of general anaesthesia and postoperative cognitive dysfunction

Reference	Age (n)	Type of surgery	Type of psychometric tests	Timing of post-operative tests	Results	Long-term change?	Comments
32	> 60 (1218 patients, plus 321 community controls)	Major non-cardiac surgery (if preoperative Mini-Mental Status was 24 or more)	EUBT (see text for details)	7 days 3 months	Postoperative cognitive dysfunction (defined after comparisons with untreated controls) in 25.8% at 7 days and 9.9% at 3 months (controls 3.4%, 2.8%)	Yes	Age, duration of anaesthesia, less education, and respiratory and other complications (but <i>not</i> hypoxaemia) correlated with dysfunction at 7 days Age was <i>only</i> risk factor correlating with cognitive dysfunction at 3 months
33	48–88 (112)	Transurethral resection of prostate	Choice reaction time	1, 2, 3 days	Increased variability in choice reaction time, day 1 only	Not tested	Increased variability associated with previous low CAPE (Clifton Assessment Procedure for the Elderly), extent of surgery, postoperative pain/sedation
16	25–83 (40)	Cholecystectomy	Mini-Mental, digit symbol/span, trail making	1, 2, 3, 4 days, 1 month	Changes on day 1 only: digit symbol (all), trail making (old only)	No	Concluded there was no major difference in rate of recovery between young and elderly groups
34	Two groups: young, mean age 50; old mean age 69 (n = 85)	Orthopaedic, gynaecological and general surgery	17 Questions (orientation/concentration), plus object learning test	2 days	Memory deficits (young and old). Orientation and concentration deficits (old only)	Not tested	Correlation between postoperative deficits and poor preoperative cognitive function

continuous pulse oximetry before surgery, throughout the day of surgery, and for the next three nights. Blood pressure was recorded by oscillometry during the operation and every 30 min for the rest of the operative day and night. Patients received general anaesthesia but no restriction was placed on anaesthetic or surgical technique, which conformed to local practice in the study centers. However, to avoid the cerebral vasoconstrictor effects of hypocapnia, capnography was a requirement during surgery, so that normocapnia could be maintained.

Analysis of the data from ISPOCD1³² showed that 25.8% patients had POCD 7 days after surgery and that 9.9% of all patients still had evidence of POCD on the repeat neuropsychological tests carried out at 3 months (corresponding values for controls were 3.4% and 2.8%). Contrary to expectations, no relationship was found between hypoxaemia and/or hypotension and the development of early or late POCD. Indeed, despite analyses of the effects of more than 25 other clinical parameters, only age showed a statistically significant correlation with late POCD. Age was also positively correlated with early POCD, as were duration of anaesthesia, a lesser level of education, a second operation, postoperative infections and respiratory complications.

The overall conclusion of the ISPOCD1 investigators³² was that their study had demonstrated a measurable degree of postoperative cognitive change in a minority of older patients 3 months after surgery (in about 10% of patients vs. about 3% of non-operated controls) and that the risk increased with age. However, the expected relationship between hypoxaemia and/or hypotension and POCD did not emerge in the study. It was also disappointing that, despite a large number of statistical analyses, the study failed to find any specific risk factors that were amenable to therapeutic or preventive intervention. In addition, the hope

that the study would give better insight into the pathophysiology of POCD was not fulfilled.

Because the ISPOCD1 study did not provide the expected answers in regard to the prevention or treatment of POCD, ISPOCD2 is now under way, coordinated from Copenhagen by J. T. Moller and L. S. Rasmussen. I am very grateful to Dr Christopher Hanning, a member of the ISPOCD2 steering committee, for the following information. ISPOCD2 comprises a linked group of multicentre projects which will ask a dozen major research questions in a variety of patient populations. A major task of ISPOCD2 will be to follow up patients for a prolonged period, to see whether the 3 month postoperative neuropsychological changes persist, and to test whether these changes produce measurable effects on Activities of Daily Living and Quality of Life. Other research tasks addressed by ISPOCD2 include: an investigation of the effects of outpatient anaesthesia; a comparison of the effects of regional anaesthesia with those of general anaesthesia; a comparison of POCD incidence in patients aged 40–60 years and in older patients; a test of the hypothesis that there might be a genetic predisposition to POCD related to the apolipoprotein E allele, which is known to have an association with the development of Alzheimer's disease; a correlation of blood levels of benzodiazepines and their metabolites with the development of POCD; an examination of the role of cholinergic and other neurotransmitters in POCD, using an animal model and positron emission tomography (PET) in humans; the study of possible protected effects of ondansetron; a study of structural cerebral changes by both MRI and SPET scanning; correlations of POCD with neurone specific enolase and protein S100; and the investigation of the relationship between POCD and "stress", particularly prolonged hypercortisolaemia.

CONCLUSION

While the great majority of older people undergoing surgery and anaesthesia will emerge without any cognitive sequelae, absolute guarantees cannot be given. Indeed, since the last edition of this book, the previous consensus that routine anaesthesia *per se* had essentially no long-term effect on postoperative function has been challenged by large carefully designed studies. It is to be hoped that the ISPOCD2 study and similar large-scale investigations will allow us to give more authoritative advice to older patients undergoing surgery in the first decade of the twenty-first century.

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Nutritional State

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Dietary surveys of elderly people in the UK and Sweden have shown that a substantial proportion of subjects had intakes below recommended standards. Two US National Health and Nutrition surveys showed that 50% of the population, especially the elderly, were deficient in one or more nutrients. In 1979, a Department of Health and Social Security (DHSS) survey in the UK suggested that 7% of those aged 65+ may be undernourished and twice this proportion of those aged 80+¹. A survey in Sydney found that intakes of nutrients by old people in Australia were similar to those recorded in the UK² and concluded that a significant proportion of this population may be at risk. Depending on definitions and diagnostic criteria, undernutrition in community-based elderly people is in the range 0–82% and in hospital 5–63%³.

High-risk groups include the housebound and those with impaired cognitive function, a history of depression, chronic pulmonary disease and partial gastrectomy. Poor dentition, swallowing difficulties and not having regular cooked meals are further factors placing people at greater risk of malnutrition.

The reasons for poor nutrition are various and the cause often multifactorial. Nutritional deficiency rarely occurs in isolation but is usually consequent upon ill-health or poverty. The UK DHSS survey of 1968⁴ found 3% of 1000 elderly people to be malnourished, largely because of untreated underlying medical conditions or socioeconomic disadvantage. Poor financial reserves, isolation, physical handicap, dental problems and mental disorder all contribute, and the effects of disease, disability and ageing can combine to change a marginally sufficient diet into a grossly deficient one⁵. The prevalence of malnutrition is certainly higher in old people than in younger groups, attributable in the main to higher proportions of the elderly living in poverty and a greater prevalence of disease in old age. The diet of lower socioeconomic groups provides cheap energy and is lower in essential nutrients, such as calcium, iron, magnesium, folate, other B vitamins and vitamin C⁶.

Methods of food preparations and a lack of knowledge about daily requirements are also important. It must be cause for concern that elderly people in residential care and long-stay hospital wards have been found to be at greatest risk of undernutrition, although naturally, they represent some of society's oldest and most frail individuals. Burns *et al.*⁷ found a group of elderly dementing people in institutional care to be more undernourished than those living alone in the community, even though they were no more cognitively impaired. Kennedy and Henderson⁸ found the risk of nutritional deficiency to be greater in residential than in hospital care.

Classical single deficiency states are rare in the UK and USA elderly, but there is a strong suspicion that suboptimal nutrition

frequently contributes to ill-health in old age. Malnutrition leads to lowered physical strength, greater inactivity and risk of accidents, a weaker immune system and osteoporosis⁶. Poor vision, macular degeneration and cataracts are being linked to diets lower in fruit, vegetables and antioxidants⁹. Subclinical vitamin deficiency is more common in the elderly, particularly those with psychiatric disorders. Hancock *et al.*¹⁰ compared nutritional indices of healthy and mentally ill elderly women and reported lower levels of vitamin C, riboflavine and pyridoxine among the elderly mentally ill. Of 255 psychogeriatric admissions with organic and functional disorders, Bober¹¹ found low concentrations of ascorbic acid in over 50% and low plasma folate in approximately 20% of subjects.

A study in England conducted by the Health Advisory Service 2000, commissioned by the Secretary of State for Health, identified feeding and nutrition in acute hospital wards for the elderly to be deficient in many respects¹². Malnutrition is largely unrecognized in British hospitals and is particularly common in old age¹³. In one study of adult hospital admissions, the overall average weight loss was 5.4% and was greatest in those initially most undernourished. Nutritional status was infrequently recorded¹⁴.

There are difficulties in the assessment of nutritional status and establishing accepted recommendations for daily requirements. Commonly used reference data in the UK may not be appropriate for all populations of elderly people, and contemporary reference data that are widely applicable, with clinically defined criteria for undernutrition, are needed¹⁵. Furthermore, it is not clear how illness may alter these requirements. The assessment of nutritional status is complex and a single measure usually insufficient; however, anthropometric measurements are the preferred clinical tool for routine use and can provide practical and valid indices of nutritional status¹⁶. Preferably, evidence of undernutrition is established from a combination of history and examination, anthropometry, haematological and biochemical indices^{17,18}. Of the 10 risk factors described by Davies¹⁹, depression, loneliness and alcoholism feature, in addition to social variables and factors reflecting dietary intake and weight change. Bender²⁰ points out that tables of nutrient requirements are of limited value, as individuals' energy and nutritional needs vary.

Although discussions of nutrition tend to focus on deficiencies, it should be remembered that the commonest nutritional disorder affecting old people in developed countries is obesity. The problem of obesity, which is often difficult to remedy, has established implications for physical illnesses such as hypertension, heart disease, diabetes mellitus and arthritis. Drugs used in the treatment of mental illness in old age not infrequently lead to weight gain.

Low body weight is associated with an increased mortality of elderly people living at home²¹ and admitted to hospital²². Poor nutrition increases the risk of falls and hip fractures²³ and is associated with excess winter deaths²⁴, the development of pressure sores²⁵ and infections²⁶.

Undernutrition may arise from quantitative and qualitative dietary inadequacy, leading to mixed nutrient deficiency. If old people meet their energy requirements by taking a good mixed diet their needs for nutrients will be met. Some elderly people lead such sedentary lives that their energy requirement is very low²⁷ and they become at risk of taking inadequate protein, minerals and vitamins. As life becomes more sedentary and energy intake declines, then diet needs to become more nutrient-dense. Asplund *et al.*²⁸ found that 30% of 91 patients, with a variety of diagnoses, admitted to psychogeriatric inpatient care were undernourished. A high prevalence of thiamine deficiency has been reported, especially among the housebound, solitary and confused²⁹ but also among admissions to psychiatric units³⁰. Alcoholism is commonly associated with deficiencies in thiamine but also riboflavine, pyridoxine, vitamin B₁₂ and folic acid. Subnormal levels of vitamin C, thiamine and pyridoxine were found by MacLennan *et al.*³¹ in long-stay elderly patients who were also protein-deficient. Inadequate intake of vitamin C has been shown for 75% of elderly men and over 80% of women, not always reflected by leucocyte ascorbate levels or laboratory parameters^{31,32}. Vitamin D levels have been shown to be very low in long-stay elderly populations³³, almost certainly secondary to inadequate exposure to sunlight³⁴. Evidence of multiple vitamin deficiencies has been reported in elderly day hospital attenders and residents of local authority residential homes⁸.

Vitamin deficiency is rarely sought in clinical practice, when only B₁₂ and folate estimations tend to be performed with any regularity. Yet any deficiency, particularly involving the B group, can present with apathy, anorexia, weight loss, mood changes, acute and chronic confusional states and occasionally hallucinations and paranoia³⁵. Depression and alcoholism seem to be the characteristic disorders associated with vitamin B and folate deficiencies, while organic psychosyndromes are typical of B₁₂ deficiency³⁰. Thiamine deficiency can produce a wide variety of mental disturbance and ascorbic acid deficiency is usually linked with apathy and depression. Irritability, aggressive behaviour and personality change were reported in healthy volunteers undergoing thiamine restriction³⁶. Vitamins and minerals are intimately involved in cell metabolism, neurotransmitter synthesis and cell membrane stability. Mineral and electrolyte deficiencies, including iron, calcium, potassium and magnesium, are also common findings among elderly populations³, with obvious implications for health.

TREATMENT

The immediate significance of single deficiency states is confined to few specific circumstances, e.g. thiamine and the Wernicke-Korsakoff syndrome, B₁₂ and folate with certain dementias and pseudodepressive states. In most instances, the import of nutritional status is less obvious and the effects of deficiency probably more subtle. Although nutritional supplementation is unlikely to be curative in these situations, there are few studies of the effects of nutritional intervention in elderly populations with mental disorder and more information is needed before drawing firm conclusions. While the real significance of nutritional manipulation is awaited, a pragmatic position recommending dietary supplementation and adjustment with efforts to prevent undernutrition is advised.

In many circumstances, treating an underlying mental disorder or physical illness effectively will restore appetite and drive,

thereby correcting deficiency by the resumption of a normal diet. The admission of a confused elderly person to hospital or care may provide the opportunity to re-establish a normal eating pattern. In specific deficiency disorders the prescription of necessary supplements will be an essential part of treatment. The possibility that nutrient supplementation may enhance the response to conventional psychotropic medication³⁷ is an interesting possibility that requires further exploration.

The correction of deficiency does not necessarily involve prescribed medication but may be possible by simple measures, such as supplementing the diet with fruit juices to provide more vitamin C. Low levels of vitamin D are found in up to 40% of the elderly living at home or in hospital. Diet is an inadequate source of vitamin D, which depends on exposure to ultraviolet light for its formation. A greater exposure to natural sunlight is the most important preventive measure, but because the elderly are at special risk it is recommended they receive vitamin D supplementation to achieve a daily intake of 10 µg³⁸.

The overwhelming priority in the management of undernutrition among the elderly population is prevention. A major impetus must ultimately come from a change in public policy that improves the elderly person's social, material and financial position in society and ensures the efficient provision of services to those in need.

The market-led approach to nutrition that operates in many food-rich countries has been found to increase the disparity between the nutrition and health of the rich and poor³⁹. The provision of domiciliary care services is inequitably distributed, often inefficiently organized and frequently determined by demand rather than need⁴⁰. Consequently, invaluable services, like meals-on-wheels and home helps (often the only people to provide food to isolated mentally ill old people), may not reach those most at risk. Little attention or imagination has been given to the meals service, potentially an important resource, which suffers from a complex, multi-agency organization, inflexibility in delivering meals of a type or at a time to suit individuals and often arriving cold. Only half the recipients find the meals at least moderately satisfactory⁴¹ while 15% never eat them and 15% eat only half those delivered⁴². There is little evidence that they are targeted at those most at risk of undernutrition and some evidence that the meals, themselves, are nutritionally inadequate⁴⁰. This has led some to suggest that the service acts only as a symbol of concern for elderly people⁴³.

Often the supervision of meal times is as important as the meal itself. Confused elderly people may eat voraciously in the company of their family yet put meals aside when left to eat alone. Altering the timing of home help or family visits or attendance at a day centre or luncheon club may be the intervention required.

The prevention of undernutrition may be possible by simple measures, e.g. providing meals in a form that is appealing and easily edible. Kennedy and Henderson⁸ demonstrated the importance of noting the food returned after meals by the elderly in residential care. Fruit, vegetables and meat were often left because they were difficult to chew. For seriously impaired individuals, maximizing food intake during the times of day when cognitive abilities are at their peak can improve dietary intake⁴⁴. The time allowed for meals is normally less in institutions than at home⁴⁵, meal times are inflexible and little consideration is given to personal choice. Some patients will take food from family members and not care staff, or only from certain members of staff. Obviously adequate staffing levels will affect success when large numbers of disabled or resistive patients are eating together.

The recent practice in UK hospitals of extending menus to include less traditional dishes may be appreciated by the young, but experience suggests many elderly people are happier with local dishes and attempting radical alterations of lifelong dietary habits

can be difficult and may be counterproductive. Changes of this sort may actually reduce the opportunity to provide a nutritious diet to elderly hospitalized patients at a time when it is most needed. Fresh fruit is often difficult to obtain in institutions and modern methods of mass food preparation have been criticized. It may not be appreciated that less mobile elderly people have a fast protein flux that requires a higher, not lower, daily requirement for protein^{17,18} and the protein intake of long-stay elderly patients can be inadequate³¹.

Poor dentition is associated with undernutrition⁴⁶ and the fitting of dentures to old people in institutional care has been shown to increase the consumption of raw vegetables⁴⁷; 78% of independent elderly patients examined by MacEntee *et al.*⁴⁸ believed their oral health to be good but only 17% had clinically healthy mouths. A study of hospitalized patients aged 61–99 years found that 60% had disease of the oral soft tissue⁴⁹.

Medication can impair the absorption and reduce the availability of essential nutrients, impair appetite, cause dry mouths and constipation or directly promote the loss of minerals, as with diuretics and potassium⁵⁰.

For those living in the community, preventive health care with early recognition and treatment of illness is essential. The elderly population may benefit from greater education and advice about healthy and affordable eating, issues normally targeted at younger age groups. The judicious use of fruit juices, frozen vegetables and some convenience foods might ease the burden of food preparation. Realizing that visual impairment and arthritic joints can prevent shopping and the opening of packets and tins may point the way to practical interventions, such as the provision of domestic aides or arranging for someone to collect shopping. One survey found that 22% of elderly people who had difficulty opening screw-top jars had to ask non-household members to do it for them⁵¹. The teaching of culinary skills is particularly relevant to the older bereaved male who never cooked when his wife was alive. Men, although age-for-age fitter than women, are twice as likely to say they are unable to cook a main meal⁵².

In the modern era shops themselves are often large, impersonal, confusing places and sited some distance from home, making effective shopping difficult for physically and mentally disabled people. Low income and disability not only restricts ability to afford a protective diet but also limits access to retailers where healthy food can be purchased more cheaply. Local shops are less prevalent and can be significantly more expensive than more distantly sited supermarkets⁵³.

Finally, unless people are aware of the possibility of undernutrition and able to make an assessment, little progress will be made for the sick and vulnerable. Sadly, doctors and nurses frequently fail to recognize undernourishment because they are not trained to look for it⁵⁴ and medical students' knowledge of the issue of nutrition is poor⁵⁵. Improved education is greatly needed. The Royal College of Nursing⁵⁶ provides clear guidelines for the assessment of nutritional status in older people.

CONCLUSION

A great deal needs to be known about the fundamentals of diet, nutrition and mental health in old age. The evidence connecting nutrition and morbidity suggests this is an area of importance to all professionals working with elderly people and a strong case could be made for the regular involvement of a dietician in the psychogeriatric team.

Establishing roles for nutritional intervention offers prospects of simple and economic measures that may improve the treatment and prognosis of mental disorders in old age, enhance clinical recovery and reduce morbidity.

At the present time, it cannot be claimed that vigorous dietary intervention offers curative treatment for mental illness but attention to diet may, at least, reduce the physical complications of mental disorder, hospitalization and ageing. However, there is accumulating interest in the role of antioxidants in the treatment of dementia [see Nutritional Factors in Dementia] and the possible significance of omega polyunsaturated fatty acids for maintaining the development and integrity of neuronal function has obvious relevance to severe mental illness⁵⁷. Schizophrenic patients taking additional omega 3 polyunsaturated fatty acids may experience milder symptoms⁵⁸ and a recent study of community residing schizophrenic patients aged 20–79 years demonstrated nutritional deficiency, despite most being overweight, with a high intake of saturated fat and low intake of antioxidant⁵⁹.

Further exploration of the relationship between dietary constituents and the course of mental illness may yet yield information significant to the management of mental illness in old age.

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Mental Illness in Nursing Homes and Hostels in Australia

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Australia has a population of over 19 000 000 people. Less than 2% are descended from the original inhabitants and over 12% are aged 65+. The probability of using nursing home care at some point in one's life is currently estimated to be around 25% at birth, rising to 60% from when aged 80 and 95% from when aged 90. As the life expectancy of those aged 65 now extends well beyond 80 years, the majority of older Australians can expect to have some experience of residential care before their lives end¹. In recent years Australian government policy has directed funding away from residential provision towards community care, and has

acted to blur the distinction between nursing homes providing full nursing care 24 h/day and hostels that offer accommodation, meals, supervision and some assistance with activities of daily living but do not furnish residents with 24 h/day nursing care^{2,3}.

In 1993 there were 74 494 nursing home beds in Australia and by 2011, if the intended planning ratio of 40 beds/1000 persons aged 70+ has been reached, there will be 78 600 beds⁴. The policy direction for hostels is to increase the level of supply from a 1993 level of 40 places/1000 persons aged 70+ to a projected level of 52.5 by 2011. There were 54 429 hostel places in Australia in 1993.

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By early 1999, 140 000 nursing home and hostel places were available in Australia, with 2500 new places being established each year, mostly in hostels¹. These places serve 2 154 000 Australians aged 65+, of whom 129 600 were thought to have dementia in 1995⁵.

Few residential facilities offer specialized care for the elderly with mental illness. In the State of Victoria a small number of nursing homes subsidized by the State Government, called "Psychogeriatric Nursing Homes", offer specialized care to elderly people with mental health problems (usually dementia), associated with behaviour too challenging to be managed in mainstream facilities. Long-term care in large mental hospitals and state geriatric centres is virtually a thing of the past⁶.

A small number of researchers have examined the prevalence of psychiatric disorders among elderly people in Australian nursing homes and hostels. At least 50% and possibly 80% of nursing home residents have dementia⁵⁻⁷. Around 40% of hostel residents have cognitive impairment consistent with dementia^{6,8}. Although the more severely demented residents cannot be assessed, it is unusual for an assessable resident to have no symptoms of depression at all and at least 10% suffer a depressive disorder at any time^{9,10}. No detailed statistics are available for the numbers of individuals with schizophrenia and related disorders living in nursing homes in Australia. The four specialist psychogeriatric nursing homes in the author's own catchment area, which serve a population of over 120 000 elderly, have 120 beds, of which around 30 are occupied by individuals with a primary diagnosis of schizophrenia and related disorder. Most of these residents are former long-term inmates of psychiatric facilities, so this percentage is likely to fall in future as fewer individuals with schizophrenia will experience long-term incarceration. The percentage of individuals with schizophrenia and related disorders in ordinary nursing homes and hostels would be far lower than this.

Despite high levels of depression and dementia, a recent study by Reberger, Hall and Criddle¹¹ revealed that entering a hostel can lead to an overall improvement in quality of life, although the size of the study was small, assessing only 50 subjects.

General practitioners are responsible for the medical care of the vast majority of individuals in nursing homes and hostels, as few residents see a psychiatrist once, let alone on a regular basis. A well-conducted study of over 2000 residents in 46 Sydney nursing homes revealed very high levels of psychotropic drug prescribing to this population¹². Psychotropic drugs were taken regularly by 58.9% of residents and another 7% were prescribed such drugs on an "as-required" basis. Antipsychotic drugs were taken regularly by 27.4% and on an "as-required" basis by a further 1.4%. These drugs were more likely to be given to residents with greater cognitive impairment and more disturbed behaviour. Benzodiazepines were prescribed to 32.3%, hypnotics to 26.6% and antidepressants to 15.6%. At least half the antidepressant doses were subtherapeutic.

As in other countries, one major concern has been the underdetection and undertreatment of depression in residential care. An innovative and painstaking series of studies were done in North Sydney by Llewellyn-Jones *et al.*¹³. This project included a randomized, controlled trial, with control and intervention groups studied sequentially and blind follow-up after 9.5 months of 220 depressed residents in a large residential facility. The interventions consisted of multidisciplinary consultation and collaboration, training of general practitioners and carers in the detection and

management of depression, and depression-related health education and activity programmes for residents. The control group received routine care. There was significantly more movement to less depressed levels of depression, as measured by the Geriatric Depression Scale (GDS) at follow-up, in the intervention than the control group. Multiple linear regression analysis found a significant intervention effect after controlling for possible confounders, intervention groups showing an average improvement of 1.87 points on the GDS. Although the impact of this study on total GDS scores was not huge, small movements in depressive symptomatology in populations are likely to be associated with significant decrease in morbidity among some individuals. In the past it has been hard to show that intervention programmes in these populations can be efficacious¹⁰, but the work of Llewellyn-Jones' team suggests that the future may not be as bleak as some of us had feared.

There is no doubt that mental illness is common among individuals who live in residential care in Australia. The challenge for our health professionals is to improve the detection and management of these conditions. A multifaceted approach is required, with improved medical education for both undergraduates and general practitioners, education for care staff and an overall improvement in the quality of residential provision for older people. Slow, relative economic decline in Australia⁶, which continues apace, will make this a difficult challenge to rise to, but Australia's track record in this area suggests that the goal is not an unachievable one.

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Caregivers and Their Support

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As this book clearly demonstrates, Alzheimer's disease and related dementias (ADRD) are tragic, debilitating, chronic illnesses with an unpredictable clinical course. The tragedy is compounded by the consequences associated with caregiving, such as impaired mental and physical health, and disruption of normal activities and social relationships of the caregiver. Cognitively impaired elders are often placed in long-term care facilities when family caregivers are either no longer available or are unable to continue home caregiving because of inability to manage behavioral problems, unremitting stress or caregiver illness. This chapter highlights the importance of the caregiver role, as well as costs to, and support for, family caregivers in the community and throughout the process of relocating the recipient to an institutional setting. Research on caregiver stress and the positive and negative effects of caregiving is reviewed, with attention to both psychosocial and physiological outcomes, as well as variables that may moderate stressors inherent in the caregiving role.

IMPORTANCE OF THE CAREGIVER ROLE

Most older adults requiring assistance reside in the community, including those with dementia¹. The main reason why persons with dementia are able to reside in the community is because they receive care from their families². Of the more than 4 million Americans with ADRD, over 70% are cared for by family members, and spouses typically are the primary caregivers³.

Families play a paramount role in the provision of care to persons with dementia and most families want to retain the role of caregiver, since their efforts can forestall institutionalization⁴. Consequently, dementia caregiving has emerged as one of the most serious healthcare issues facing our society today⁵.

Who Are Family Caregivers?

There is no precise definition of family caregiving to guide researchers and health professionals^{6,7}. Caregiving is a truly heterogeneous concept. Family members define caregiving and care-receiving differently, depending upon their relationship to the care recipient, gender, and whether or not residence is with the care recipient.

Informal caregiving typically falls to women, who provide 72% of family caregiving, with adult daughters providing 29% and wives 43%⁸; 80% of caregivers of persons with dementia are

women, 55% are spouses, 35% are adult offspring and 5% are siblings; the remainder are other relatives or paid providers. Thus, family caregiving is more often a career for women⁹. Wives and daughters provide more activities of daily living (ADL) and instrumental activities of daily living (IADL) assistance to family members than do husbands and sons^{10,11}. The preponderance of customary IADL assistance, viewed as "women's work", may result in the underestimation of the amount of caregiving of persons with dementia that is borne by women. However, there is evidence that men and women caregivers do not differ in length of caregiving service, hours per day spent in caring for relatives, and the perceived stress of caring for persons with dementia. Both perceive the most stress from dealing with behavioral problems, a common precipitant of institutionalization¹².

Public Policy and Cost Implications of Caregiving

Factors such as the aging of the population, the increase in dependency with age, and the scope of care provided by family members are especially important in understanding dementia care. Policy makers are increasingly aware of the fact that family caregiving is not cost-free. Caregivers incur numerous expenses, including home modifications, assistive devices, special food, high utility costs, and the cost of foregoing paid employment. This type of family support is estimated to be the equivalent of full-time work in about one-third of households providing dementia care. According to Day¹³, the cash value of services performed by family caregivers far exceeds the combined cost of government and professional services to both the elderly who live in the community and those who live in institutions. Current estimates indicate that dementia family caregivers save the US healthcare system \$196 billion/year¹⁴. Therefore, the importance of family caregivers will increase in the future, as the number of persons with dementia, and the costs of their care, increase.

Society at large benefits from the willingness and ability of families to care for their members who have dementia. Were it otherwise, persons with dementia would be institutionalized sooner and in larger numbers than is currently the case. This, in turn, would impose a far greater economic burden on society in providing long-term care to persons with dementia. Importantly, monetary estimates of dementia care in no way reflect the human costs of this devastating disease. Although family caregiving helps to contain costs to society as a whole, it often results in serious costs that are concentrated in caregiving families. As noted, the breadth and magnitude of these costs cannot be

calculated solely in economic terms. A large body of research has established that there are also profound effects of caregiving associated with poor health-related outcomes^{15,16}, as detailed later in this chapter.

CAREGIVER STRESS AND BURDEN AND ITS IMPACT ON CAREGIVER WELL-BEING

Concomitant with an increase in the prevalence of dementia is an escalation of the physical, emotional and economic burdens of dementia care. The concept of caregiver burden has been advanced as an all-encompassing term that refers to the financial, social, physical and emotional effects of caring for a family member with dementia¹⁷. Subjective burden, the caregiver's perception of the caregiving experience, is distinguished from objective burden, the actual changes in the caregiver's home situation¹⁸. Studies of the burden and stress of caring for a person with dementia, the resulting effects on mental and physical health outcomes of family caregivers, and the use of a variety of interventions to relieve adverse effects of caregiving burden are numerous. More recent research also has focused on the positive aspects associated with caregiving. Major findings from this body of research are summarized throughout this chapter.

Family caregivers of persons with ADRD experience numerous stressors that affect their health and well-being and precipitate institutionalization of the care recipient. Hence, the notion of "stress" has emerged as an important concept in dementia caregiver research³. The words "stress", "burden" and "distress" are often used interchangeably and are conceptualized to have similar meaning. George and Gwyther¹⁹ defined the concept of dementia caregiver stress as: "the physical, psychological or emotional, social and financial problems that can be experienced by family members caring for impaired older adults" (p.243). A variety of factors related to the symptoms associated with ADRD cause caregiver stress, including cognitive changes, loss of ability to function in daily activities, and behavioral disturbances. Spouses of persons with dementia may be at greatest risk for caregiver stress, since they are often elderly themselves. They may also have physical, psychological and financial challenges that could decrease their ability to respond to the demands of caregiving^{20,21}.

Models of Caregiver Stress

A number of theoretical frameworks have been used to guide studies of persons with dementia, their family caregivers and the effects of interventions on outcomes for both²²⁻²⁸. Stress models, in particular, have guided much of this research. However, models that predict a simple positive linear relationship between caregiver stress and the care recipient's level of impairment, such that caregivers providing assistance to the most behaviorally impaired persons report the greatest degree of burden, are not supported by the literature^{29,30}. Rather, research indicates that caregiver stress is related to a number of care recipient and caregiver variables³¹ and, as such, is a multivariate phenomenon.

Caregiver stress is moderated by many factors, including the caregiver's available resources, such as good physical health, social support, financial assets, coping abilities and personality

$$\text{Distress} = \frac{\text{Exposure to stressors} + \text{Vulnerability}}{\text{Psychological resources} + \text{Social resources}}$$

Figure 138b.1

traits¹⁷. Although a number of multivariate models of caregiver distress exist, space limitations preclude a review of extant models in this chapter. Rather, Vitaliano's model of distress³² (Figure 138b.1) is set forth as an example of one useful model for understanding caregiver stress, because the variables of interest are well grounded in the caregiving and theoretical stress literature. Additionally, the model allows for the stratification of resources and vulnerability variables, which improves the chances of detecting relationships among stressors, resources and burden. In Vitaliano's model, both caregiver and care recipient variables, as well as psychological and social resources of the caregiver, are postulated to contribute to caregiver distress³².

The underlying assumption of research on caregiving stress and health outcomes is that the chronic stress of caregiving can lead, via vulnerabilities and limited resources, to psychological or physiological distress and illness. Vitaliano's model, for example, considers both psychological and biological markers of distress. Since data overwhelmingly suggest negative psychological consequences of caregiver stress, biological outcomes allow for assessment of the impact of the psychological distress on major physiological systems. As noted earlier, in caregiver research, multivariate models provide a more comprehensive picture of a caregiver's level of distress than either psychological or physiological variables alone³². Moreover, since Vitaliano's model of distress is expressed as a mathematical formula, a caregiver's level of distress (burden) may improve by decreasing undesirable variables or by increasing desirable variables.

STRESSORS ASSOCIATED WITH FAMILY CAREGIVING

Providing care for a family member with dementia is conceptualized as a chronic stressor³³. Symptoms of dementia that the caregiver must contend with include (but are not limited to) progressive loss of: memory; judgment; the ability to interpret abstractions; language and motor deficits; and altered personality. Dementia results in profound cognitive and behavioral changes that culminate in an inability to perform instrumental activities of daily living, such as cooking and managing money, as well as basic activities of daily living (ADLs), such as bathing and toileting. The level of functional (ADL and/or IADL) impairment in care recipients was related to caregiver burden in one study¹⁵, although a larger number of studies found no association between these variables³⁴⁻⁴¹. Similarly, no evidence of a relationship exists between the care recipient's level of cognitive impairment and the caregiver's level of burden^{34,35,39,42-44}.

The course of the illness is unpredictable, as is the rapidity of decline; the only certainty is that the progressive cognitive impairments that characterize dementia will lead to an increasing need for supportive care and eventual death for the person with dementia. As the illness progresses, caregivers must be increasingly vigilant, since dementia patients may elope from home or injure themselves. In the final stages, patients are often completely dependent on their caregivers and need to be fed, bathed, toileted, transferred and dressed. Providing care to a person with dementia eventually becomes an all-consuming 24-h job, which may extend for 10 years or longer^{45,46}.

Behavioral Impairments in Care Recipients

The stress of providing 24 h care for a person with dementia is complicated by the development of episodic, problematic (or catastrophic) behaviors, when the person with dementia

becomes increasingly agitated, stressed or disorientated. The most commonly reported behavioral changes associated with dementia include neurovegetative symptoms (e.g. lethargy, social withdrawal), sleep disturbance, restlessness, wandering, assault, aggression, destroying property, verbally disruptive behavior (e.g. screaming), and inappropriate sexual behavior⁴⁷⁻⁵¹. Behavioral problems are extensive in persons suffering from dementia, appearing in up to 67% of care recipients upon diagnosis⁵², approximately 65% of demented persons who are institutionalized⁴⁹, and 70-90% of persons with advanced dementia^{53,54}. They worsen with disease progression and may be related to fatigue, change, overstimulation, excessive demands or physical stressors²³. Problematic behaviors appear to have a profound effect on caregivers' stress, and a number of investigators have concluded that these secondary symptoms are the most stressful to manage from a caregiving perspective^{46,55,56}. In fact, the one care recipient characteristic that overwhelmingly predicts caregiver distress is the degree to which the care recipient demonstrates behavioral problems^{15,34,37,44,47,50,51,57}. Many investigators also report that behavioral problems are strong predictors of institutionalization^{47,57,58}.

Other Factors that Influence Caregiver Stress

As noted earlier, researchers typically theorize the stress of caregiving as chronic because caregivers face many years of continuous exposure to the daily demands of caregiving. However, over the disease trajectory, the intensity and/or frequency of a caregiver's level of distress may vary widely. In an effort to better understand this variability, investigators have identified a number of factors associated with differing levels of morbid outcomes. Of special interest are a number of variables that appear to moderate, or render less harmful, the effects of caregiving among some individuals.

It appears that the amount of stress is influenced by whether or not the caregiver is co-resident with the care recipient, the abruptness of onset of the care recipient's disease, kinship relationship with the care recipient, and the coping strategies used by caregivers^{59,60}. Seltzer and Li⁶¹ found that daughters in later stages of caregiving had a more distant relationship with the care recipient and more subjective burden than daughters in the earlier stages, while wives evidenced the opposite pattern. Wives who had provided care for a longer time reported less burden and a closer relationship with their husbands if they perceived themselves to be in the later stages of caregiving. This finding is supported by the longitudinal studies of Schulz and Williamson⁴⁰, which suggest that caregivers have successfully adjusted to the rigors of caregiving and have learned to cope with the demands of the task. Evidence is mixed as to gender differences in the stress experienced once the caregiving role is undertaken⁶². Overall, the literature suggests that stress-related gender differences are more pronounced for caregivers of non-demented persons¹².

Social support has been examined both as a correlate of distress (a main effect) and as a modifier of the relationship between stressful experiences and distress (an interaction effect). In the broader stress and coping literature, social support has been a consistent moderator of stress-related outcomes⁶³, in that the presence of a strong social network and satisfaction with support is a powerful predictor of positive outcomes. Caregiving studies suggest a direct effect of social support on measures of burden, but evidence for a buffering effect has been less clear³².

In sum, research has identified potential exposure (care recipient behaviors), vulnerability (e.g. age, gender, neuroticism, pre-existing hypertension) and resource (social support) variables

that may either moderate or mediate relationships of caregiving psychological distress with measures of physiological impairment and illness. In spite of overwhelming evidence that caregiving is stressful, factors contributing to caregiver distress have not yet been delineated in a way that can effectively direct interventions or preventive strategies.

Dementia vs. Non-dementia Caregiving

There is evidence that family caring for a person with dementia is more stressful than caring for a person who is not demented or who has a physical limitation⁶⁴. For example, Clipp and George⁶⁵ found that family members caring for a person with a dementia were more adversely affected by their role than family members caring for a relative with cancer. This relationship was not explained by the duration of the illness or by whether or not the caregiver was employed; however, younger spouse caregivers were more adversely affected than older caregivers. In a review of studies of caregiving in different types of illnesses, Biegel *et al.*⁶⁶ observed different patterns of distress. They concluded that the pattern of a peak distress period after initial diagnosis of acute onset, followed by a reduction in distress as time passes, was not observed in family caregivers of persons with gradual onset illness where no relief of distress was observed⁶⁷. Co-resident caregivers of persons with stroke and of persons with dementia experienced similar degrees of burden and high levels of psychological distress, with psychiatric aspects of care resulting in greater stress than physical aspects³⁶.

POSITIVE AND NEGATIVE OUTCOMES OF FAMILY CAREGIVING FOR PERSONS WITH DEMENTIA

Although family caregiving of persons with dementia is usually regarded as stressful and includes a variety of negative outcomes, there is growing consensus that this is not always the case⁶⁸. Some caregivers may receive satisfaction from the role^{69,70}. Caregivers can gain satisfactions, emotional uplifts, gains in self-esteem and self-efficacy, optimism, and growth and meaning from their roles⁷¹⁻⁷⁴.

Impact of Caregiving on Health of Caregivers

Findings of studies are inconsistent as to the effects of caring for persons with dementia on family caregivers' mental and physical health. Some report no changes in emotional and physical health and a decrease in depression over time, although far more report worsening of depression and physical health^{45,75-79}. Based on the findings from several studies^{44,59,60,80} Wright⁸¹ suggests that these inconsistent findings may be due to differences in coping strategies employed by caregivers.

Mental Health Outcomes

There is extensive documentation that caregivers are at risk for high levels of psychological distress (e.g. burden, stress, depression, perceived hassles). Despite gaps in the literature and differences in research methods used, caregiver studies have overwhelmingly pointed to the adverse effects that caregiving for someone with troublesome behavioral symptoms can have on the caregiver's mental health^{16,82,83}. A number of studies have also revealed negative changes due to the strains of direct care, grief associated with the deterioration of their loved ones, social isolation, and the role changes of caregiving, including care at

home, following institutionalization, and when the care recipient dies^{20,41,66,83-85}.

Depression

Depressive symptoms are among the most frequently examined mental health effects on family caregivers of persons with dementia. Several studies document a greater prevalence of depression among caregivers of persons with dementia, compared to other age and gender group norms and persons who are not caregiving^{43,86-88}.

Prevalence rates of depressive symptomatology among caregivers range from 30%⁸⁷ to 46% among community caregivers⁸⁶. Moreover, depression among caregivers is associated with intensity of their reactions to the patients' memory and behavior problems^{89,90} and to other adverse outcomes, such as increased physical burden³⁶, subjective burden⁹¹⁻⁹³, and use of psychotropic medications^{15,65}.

Depression is also noted to be greater among females than males¹⁰ and appears to increase over time among residential caregivers and decrease over time following institutionalization and bereavement^{83,87}. However, younger spouse caregivers were found to have higher levels of depression than older spouses in a study of residential and institutional caregiving⁹⁴. There were positive relationships between depression and health status and depression and days unable to work among residential caregivers, but only between physical health characteristics and health status among the institutional spouse caregivers, and no significant difference in depression between genders. Compared to non-caregiving men, male spouse caregivers have been shown to have higher levels of depression, respiratory symptoms and poorer health habits, but the groups did not differ on other measures of physical and mental health⁹⁵.

Although no subjects were clinically depressed, Wright⁸¹ found spouse caregivers of demented persons to have significantly greater dysphoric moods than a comparison group of non-caregiving spouses, with sadness of subsequently widowed spouses significantly greater than for non-widowed spouses. Widowed spouses also had poorer health outcomes over time and used fewer positive coping strategies, regardless of whether they placed their loved one in an institution.

Anxiety

Several investigators incorporated self-report measures of anxiety into their studies of depression. Mohide *et al.*⁹⁶ found that 22/23 individuals found to have significant symptoms of depression also reported significant symptoms of anxiety. Similarly, Vitaliano *et al.*³⁰ found 35.4% of their sample to have significant symptoms of anxiety. Neundorfer⁹³ reported much lower rates of both anxiety (15%) and depression (25%) among her sample of caregivers, although their scores were somewhat higher than population norms for elderly individuals.

Overall, there is strong evidence suggesting that caregivers exhibit higher levels of psychiatric symptomatology than comparison groups. Yet caution must be exercised when interpreting the generalizability of these findings, since many of the samples may be biased toward the more distressed members of the caregiving community. For example, the majority of caregivers are recruited from local chapters of the Alzheimer's Association, caregiver support groups, or through referrals by healthcare professionals. Individuals who have little difficulty with the caregiving role or who are so distressed or constrained that they are unable to participate in supportive programs or visit healthcare professionals may be underrepresented in research.

Some authors have also questioned the clinical significance of psychiatric symptoms reported in caregiver distress studies. There remains a criterion problem of distinguishing normal distress from psychiatric illness⁸⁴. Transient periods of grief, despair, helplessness and hopelessness may be much more common than a diagnosable depression among family caregivers. Becker and Morissey⁸⁴ argue that the severe and chronic stressors associated with dementia caregiving are unlikely to precipitate a major depressive disorder, except in predisposed individuals. They suggest that the depressive-like symptoms among caregiving spouses should be categorized under Code V: Conditions Not Due to a Psychiatric Disorder⁸⁴. This view is consistent with the conclusion reached by Fitting *et al.*⁹⁷ who state: "It is our impression that most caregivers reporting depressive symptomatology are experiencing a 'transient dysphoric mood' and not major depression" (p. 250).

To summarize, the literature on mental health outcomes of caregiving is suggestive but not conclusive⁹⁸. There is strong evidence for increased symptom reports for depression, anxiety and increased psychotropic drug use among caregivers in these studies, as well as support for increased clinical psychiatric illness among some caregivers.

Physical Health Outcomes

Most of the literature examining the physical effects of caregiving has used one or more indicators of caregiver health: (a) self-reported health status; (b) self-reported incidence of illness-related symptoms; (c) self-reported utilization of health-care services; (d) self-reported medication use; and (e) biological indicators as a measure of susceptibility to disease. Reported predictors of poor physical health outcomes for caregivers include older age, being a spouse rather than adult child or other relative, and being female rather than male^{15,79}. Interestingly, a relationship between the positive aspects of caregiving and physical health has also been reported. Emotional uplifts in family members with coronary heart disease (CHD), who are caring for persons with dementia, mediates the severity of metabolic signs that predict CHD⁹⁹.

Although there are fewer reported studies, the psychoneurological and immunological effects of caregiving are receiving increased attention. A decrease in measures of cellular immunity, and more days of infection following long-term residential caregiving of persons with dementia compared to controls also has been demonstrated, with caregivers who reported less social support and more stress having the greatest adverse immune function effects 13 months later⁸⁷. Alterations in physiological function as a result of exposure to stress have been found to increase the probability of illness¹⁰⁰.

Similarly, psychological stress can increase caregiver vulnerability to disease by compromising the integrity of the immune system¹⁰¹. Kiecolt-Glaser *et al.*¹⁰¹ have examined depression and distress as immunological modifiers. Their research team has also documented poorer immune response, in particular changes in the percentages of helper T-lymphocytes and natural killer (NK) cells, in caregivers of persons with dementia, while controlling for nutritional intake and illness-related variables¹⁰¹.

Caregivers in this study also reported nearly three times as many stress-related symptoms and higher rates of psychotropic drug use than controls, especially those who were living with the person with dementia. Other studies comparing psychotropic drug use in caregiving and non-caregiving samples have also reported that caregivers use more psychotropic medications than non-caregiving controls^{15,87,102}. Reports of somatic medication use

among caregivers are less consistent, but most studies do not demonstrate a significant difference in the use of these medications between caregiving and non-caregiving samples^{42,87,103}.

In summary, the findings for physical health effects of caregiving are more equivocal than those for mental health outcomes. Although a number of investigators report significant health effects among subsets of caregivers, patterns of findings across studies are not as consistent. Evidence linking caregiving to physical health indicators, such as reported illness, physical symptoms, healthcare utilization or health-related behaviors, is generally weak¹⁵. This may be due to different definitions of health, health outcome measures, caregiving and control samples (varying levels of vulnerability and resources), care recipient samples (functional vs. behavioral impairments) and the fact that some self-report measures of physical health may primarily reflect life satisfaction¹⁰⁴.

Evaluating links between caregiver distress and health outcomes will ultimately require complex, multivariate models that are tested prospectively, over an extended period of time. Despite methodological challenges inherent in the evaluation of caregiving outcomes, data from several laboratories lend weight to the argument that chronic stressors contribute to affective disorders and may alter caregivers' sympathetic, neuroendocrine and immunological function.

FAMILY CAREGIVING: RELOCATING THE CARE RECIPIENT

Placing a relative in a nursing home is a stressful event for both the family caregiver and the patient. Caregiving places substantial burdens upon the caregiver and these burdens increase with the progression of the disease. Yet demented persons are usually placed in nursing homes only when all other avenues have been tried and other resources are exhausted¹⁰⁵⁻¹⁰⁷. Generally, the decision is postponed long past the time when more objective persons see it as appropriate¹⁰⁸. Or decisions may be crisis-driven, e.g. the care recipient wanders off, sets the stove on fire, or overdoses on pills. One reason this delay occurs is that some caregivers, especially spouses, believe that their role obliges them to sole caregiving responsibility and to never institutionalize their relative¹⁰⁹. Others may have promised, "I'll never put you in a nursing home", or "... til death do us part". Children, while more likely than spouses to rely on formal services and less enduring than spouse caregivers, nevertheless delay placement decisions because of reluctance to reverse roles and take charge of a parent's life¹¹⁰.

Overall, the literature indicates that the care recipient's extent of cognitive impairment, loss of self-care abilities, and disruptive behaviors; mediated by caregiver age, employment status, health, stress and burden, relationship with the care recipient, duration of caregiving, and support and moderation by caregiver minority status, kinship relationship with the care recipient, and use of in-home services and resources, are predictive of institutionalization. Cohen *et al.*¹⁰⁵ described seven variables that affect a caregiver's decision to institutionalize a dependent elder with dementia: use of services; enjoyment of caregiving; caregiver burden and health; caregiver rating; reaction to care receiver behavior and memory problems; and presence of troublesome behaviors. Six variables predicted actual institutionalization: caregiver health; caregiver burden; use of services; care receiver cognitive function; troublesome behaviors; and caregiver reaction to behaviors. Montgomery and Kosloski¹¹⁰ compared predictors of placement for adult child caregivers and spouse caregivers. Higher income, eligibility for Medicaid, lower morale and age of the elder were associated with placement for both groups, but other predictors of placement were different.

Notably, level of affection predicted placement for children, but not for spouses, while sense of obligation was predictive for spouses, but not for children, who were more likely to place the care recipient in a nursing home at all points in time. The probability of placement declined with time for a while, then leveled off, then rose as caregiving duration exceeded 30 months, with the probability increasing more sharply for child caregivers than for spouses.

Although the number of community services has increased to support family caregiving in the home, the extent to which these services are meeting the needs of caregivers is questioned¹¹¹⁻¹¹³. Collins *et al.*¹¹⁴ found that 40% of family caregivers who had placed their loved one with dementia in a nursing home, reported that the availability of at least one additional community service would have delayed the institutionalization. The assumptions that family members know how to provide all of the care that is needed and that they have access to the resources to assist them with provision of needed care in the home are not valid, according to current research¹¹². As a result, studies by Archbold *et al.*¹¹⁵, Brennan *et al.*¹¹⁶ and Buckwalter *et al.*¹¹⁷ have evaluated interventions to assist family caregivers in the home with skills, anticipation of decisions and role changes and access to resources.

The consideration of relocation raises the prospect of sharp role transition. For most spouses, relocation changes a longstanding pattern of living together and providing for the other. For children it can mean the restoration of a pattern in which the child is not living with, and/or is not directly responsible for, the care of the parent. Roles that were previously reversed from parent parenting child to child parenting parent are again reversed. Interviews conducted during the Family Involvement in Care research¹¹⁸ revealed that adult child caregivers of persons with dementia found the decision to put parents in a nursing home very distressing. Frequently reported comments were: "... the worst time in my life"; "... it about killed me to do it"; "... it really bothered me because I knew she would be angry with me", or "... my brothers didn't agree with me and that was a worry". While spouses reported some of these feelings, they tended to be more concerned with the loss of spouse and of the role of caring for the mate. Comments included: "... I knew I would miss him"; "... I hated thinking about not being able to take care of him"; or "... I kept thinking about how he would probably miss me and the things I do for him".

Persons with dementia placed in nursing homes may be highly resistant and fearful of the change. Given their diminished capacity for reasoning, it can be impossible to convince them that they require institutionalization. This presents a very stressful dilemma for family caregivers. It may be more difficult for child caregivers than for spouses, because of the need to reverse roles. Constant requests to be taken home are especially stressful to families. On the other hand, spouses may find it more difficult if their long-term close relationship has been affectionate and loving. In Family Involvement in Care interviews¹¹⁸, the majority of children, spouses and other relatives noted that it was very hard to actually place the relative in a nursing home. Comments were: "... I cried all the way home"; "... it was so empty at home and I felt so lonely"; "... I knew she would miss her things, so I took as much along as I could, pictures and such. She had so much and then so little, it didn't seem fair"; or "... he kept saying he wanted to go home and tried to leave with me ... it was so sad". Many of the same family members' comments, however, indicated that they also were relieved, but ambivalent: "... it was hard, but I felt like a weight had been lifted"; "... I feel guilty saying so, but I was so glad it was finally done"; "... I felt free to do some things for myself again"; or "... it was hard to do but I knew it was best for my family".

When caregivers lack the necessary skills to manage problematic behaviors effectively, care recipient behaviors often escalate, leading to increased levels of confusion and agitation^{23,103} that have been cited as the primary reasons for institutionalizing a family member with dementia^{46,119,120}. Considering the psychological and financial expense associated with placing a person with dementia in an institutional setting, interventions which help caregivers in the prevention and/or management of behavioral problems are both timely and significant. In support of this position, the 1993 *Report of the Advisory Panel on Alzheimer's Disease* recommends that emphasis be directed toward health services research that focuses on reducing the burdens of care for family members of persons with dementia.

Although our understanding of the dementias and the severe negative consequences of caregiving for this population has grown, too little research has focused on the development, implementation and evaluation of interventions, especially for family caregivers. The American Association for Geriatric Psychiatry, the Alzheimer's Association and the American Geriatrics Society, in consensus, state: "Interventions that reduce the risk of caregiver depression and improve tolerance and the capacity to care for patients in the home, including educational materials, counseling support groups, day care and respite care, are among the most promising areas for future research"¹²¹.

CONCLUSION

Caring for a person with cognitive impairment is chronically stressful. For persons with dementia, caregiving can last an average of 15 years, with the task being a demanding and often overwhelming experience⁶². This chapter has summarized the research on family caregiving of persons with dementia in the community setting. Informal caregiving, mostly by family members, of elders with chronic illnesses and disabilities has increased during the past two decades due to growth in the proportion of elders in the population, shorter hospital stays, advancing technologies that forestall mortality, and continuing higher costs of health care, with limited reimbursement of caregiving by professionals in the home. As a result, the impact of informal caring on family members quickly became a concern of health professionals and social and behavioral researchers, especially in the context of dementia caregiving. After two decades of extensive attention by the public and researchers, there is some consensus that, while there are both negative and positive effects of informal caregiving, outcomes are different depending upon characteristics of the caregiver and the care recipient. Moreover, different interventions are effective in ameliorating negative caregiver outcomes for specific caregiver and care recipient contingencies.

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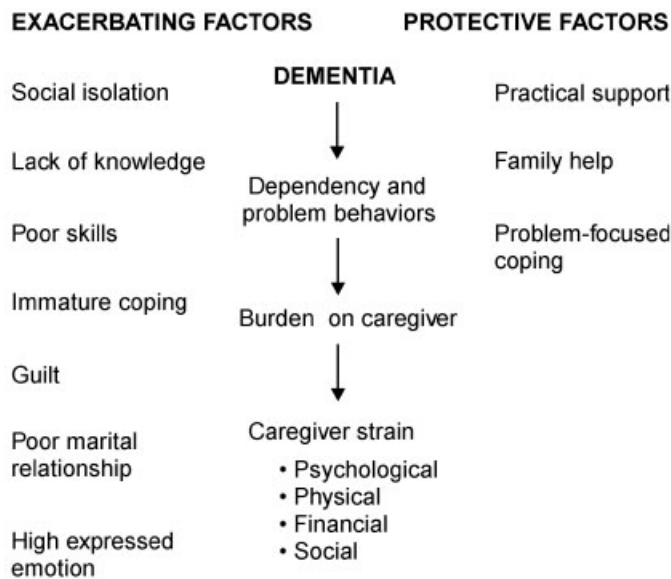


Figure 1 Model of the effects of dementia on caregivers

close to the caregiver and past history of psychological ill-health were other risk factors³⁻⁶.

By the early 1980s, several intervention techniques had been described. These were designed to enhance the skills of caregivers in coping by cognitive-behavioural approaches, training in problem solving, educational therapy, meditative relaxation, training in social skills, supportive counselling of individuals or families, and the management of stress (see reviews^{7,8}).

The outcomes of studies were limited by a number of methodological difficulties. The numbers of subjects were small, the period of follow-up was insufficient, the interventions were limited in scope, the baseline level of morbidity in subjects was often low, making it impossible to demonstrate improvement, and patient outcome was inadequately evaluated. Even so, there were some positive outcomes demonstrated: reduced family burden, decreased psychological morbidity, greater knowledge about dementia and more assertiveness and tolerance by caregivers⁷.

A number of events, some unique, led to the development of the Sydney Dementia Carers' Training Program. First was the content, which was based on addressing factors known to exacerbate caregiver distress and indirectly potentiate nursing home admission. Second, there was a period of low occupancy on the psychiatric ward of a general hospital which, if not remedied, meant that some of the beds were under threat of closure. Third, the author, as the director and superintendent of that ward, was able to utilize a variety of staff to assist in the program. Fourth, a pilot program had proved feasible and anecdotally effective and facilitated a grant from the Australian Commonwealth Department of Health.

In the model underpinning the program (Figure 1), dementia leads to increasing dependency and a number of problem behaviours. These impose a burden on the caregiver, which can manifest as psychological, physical, financial and social strain.

Exacerbating factors drive the reaction towards more psychological strain; protective factors ameliorate this.

CONTENT OF THE PROGRAM^a

The content of the 10 day intensive residential program can be conceptualized under 10 rubrics [discipline of professional(s) conducting the session; number of sessions and duration of sessions are indicated in parentheses]:

1. *Reducing caregiver distress* (social worker/occupational therapist; 2 × 2 h + 1 × 1.5 h). These sessions were scheduled first, as we found that caregivers were unable to acquire new knowledge until their psychological distress had been dealt with. Sessions were informal, supportive and expressive in nature, with caregivers encouraged to unburden themselves. Discussion in the first session included topics such as caregivers' stories, the stresses of caring, associated feelings, setting limits for the person with dementia and caregiver, coping with caring, and role changes. The second session explored the additional themes of acceptance of the disease and how dementia affects relationships with family, friends and community. The third session focused on caregiver burn-out and how to look after one's individual needs.
2. *Combating isolation* (psychiatrist; 1 × 1 h). We aimed to reduce caregivers' social isolation by the group interaction, residential setting and bringing together four caregivers for 10 days. This often led to mini-support groups forming and was a rehearsal for participation in other support groups. After the intensive residential program, telephone conference calls and hospital follow-up visits strengthened the bonds between groups of four caregivers. Extended formal family sessions brought together an expanded network of potential caregivers. For many families it was the first time they had all gathered to discuss ways of assisting with care. Sometimes geographically distant families participated on speaker telephones.
3. *Guilt and separation*. Caregivers previously trapped by their role, guilt or their partner's insecurity or suspiciousness were separated from their charges for most sessions, and encouraged to enjoy a number of activities, such as excursions to the local shopping centre or coffee shop. This provided a rehearsal for more separateness when at home.
4. *New ways of thinking*
 - (a) *Assertiveness training* (psychologist or occupational therapist; 2 × 1.5 h). Participants were provided with a working knowledge of assertive, non-assertive and aggressive behaviours, with their own "Bill of Rights", and with strategies for coping with criticism. Sessions were concrete and used role-play extensively.
 - (b) *Re-roling* (psychologist; 1 × 2 h). This focused on roles—concept, definition (mainly by gender), expectations and responsibilities—and how these were affected by dementia. Many caregivers had considerable difficulty taking over roles relinquished by the dementing person, such as driving, organizing the family and dealing with bureaucracy. Required skills were identified and their development promoted.
 - (c) *Relaxation and stress management* (occupational therapist; 8 × 30 min, daily). Techniques for relaxation, meditation, use of physical imagery and progressive muscular relaxation, complemented by two half-hour discussions on the theory of relaxation and stress response, were very popular. Caregivers obtained audiotapes to practise techniques themselves and with their partners.

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5. *New coping skills*

(a) *Communication* (psychologist; 1 × 2 h). The first half of this component was theoretical. It focused on:

- How to communicate with a dementing person.
- The functions and expectations of communication.
- How communication processes can be disrupted.
- Information on language impairment that occurs in dementia, such as receptive and expressive dysphasia.
- Techniques for clear communication, e.g. the four Ss—keep it simple, slow, short and specific⁹.

In the second half, techniques were practised by each patient-caregiver dyad while being videotaped and/or observed by the group through a one-way mirror. Caregivers were able to review the videotape and analyse their performance and communication techniques.

(b) *Reality orientation* (occupational therapist; 1 × 1 h). This was based on a 24 h environmental reality orientation model with use of verbal techniques, signs, pictures, clocks, diaries and other strategies.

(c) *The therapeutic use of activities* (occupational therapist; 1 × 1 h). This introduced the concepts of activity as being goal-directed use of time, energy and attention. *Activity analysis* was explained as breaking tasks into small steps, then modifying, eliminating or replacing steps that prevented the dementing person from completing the task (e.g. having a bath, playing golf, cooking). There was also much discussion on appropriate leisure pursuits.

(d) *Reminiscence* (occupational therapist; 1 × 1 h). Caregivers were taught how to compile a “This Is Your Life” book, comprising mementos and photographs that described the past life of the patient. This proved to be a positive experience and subsequently provided a good stimulus for conversation and reminiscing.

(e) *Coping with physical frailty* (various; 3 × 1 h). First, a physiotherapist discussed back care, walking and mobility aids. Second, an occupational therapist discussed the use and abuse of aids to daily living; caregivers tried out many of these aids in a modified kitchen, bathroom and bedroom in the occupational therapy department. Third, a registered nurse outlined the care of bed-bound, chair-bound and incontinent persons.

6. *Fitness, diet, organizing the day and home* (various; 3 × 1 h).

(a) A physiotherapist encouraged fitness and flexibility in caregivers as well as patients. For example, a daily routine of walking after lunch was established.

(b) A dietitian outlined the principles of a healthy, balanced diet and discussed time-saving kitchen techniques as well as food fads and eating problems associated with dementia.

(c) In a session on *work simplification*, and *organization and safety in the home*, the occupational therapist explored techniques of how to prioritize and simplify tasks and how and when to recruit outside assistance. The aim was to help caregivers achieve a balance between work, leisure and rest in their lives. Safety issues pertinent to the older person and the dementing process were discussed and a safety checklist for the home and garden was provided.

7. *Medical aspects of dementia* (psychiatrist; 2 × 1 h). These sessions provided medical information on dementia and its different types, principles of management, psychiatric complications and behavioural changes, use and abuse of medication, the interaction of dementia and other illnesses, and prognosis. As with all of these sessions, much time was given to answering individual concerns.8. *Using community services* (welfare officer; 1 × 1.5 h). This very practical session included procedures and eligibility for social securities, provision of useful contact persons, and

access to and availability of services. For some caregivers it was a novelty to adopt the role of *care manager*, e.g. organizing other people, such as domiciliary nurses, to help with provision of care, rather than that of *care provider*, where the caregiver undertook tasks personally. Reinforcement was given that use of services did not represent failure or dereliction of duty. Numerous pamphlets were provided on domiciliary nursing care benefits, pensions, methods of assessing nursing homes and hostels and mechanisms for complaints about services.

9. *Planning for the future* (psychiatrist; 1 × 1 h). The last formal session was fairly open and considered *how to plan for emergencies*, e.g. should something happen to the caregiver. Other issues, such as driving, medications, safe use of alcohol, smoking, legal, medical and financial matters and other emergency contingencies were discussed.10. *Coping with problem behaviours*. There was no time set to discuss these specifically, although each session was structured to allow discussion of current or potential problems, such as aggression or wandering. The aim was to give caregivers a broad education on the possible reasons for the emergence of problems and a repertoire of skills to prevent their occurrence or to deal with them if they occurred.

THE PATIENT PROGRAM

For caregivers to be able to learn in a relaxed setting, they needed to know that their partners were receiving satisfactory care. Patients had their own program, which consisted of: (a) general ward activities, such as occupational therapy, outings and relaxation classes; and (b) specific programs—group discussion of their frustrations with their memory loss, reminiscence therapy and a memory retraining program. Given a forum for honest and open discussion, patients established strong bonds with each other, were able to discuss their feelings surprisingly frankly and often became protective of each other. Memory techniques included use of visualization and one-tracking (focusing on one task to be remembered at a time). While we were unable to demonstrate any improvement in cognitive function¹⁰, our impression was that patient morale improved.

Process

The course was residential and for a variety of (non-essential) reasons, as explained above, took place in the psychiatric ward of a general teaching hospital, with caregiver and patient couples sharing individual rooms. An advantage of this setting was the availability of facilities and staff. A disadvantage was the inappropriateness of some interactions with psychiatric patients, yet there was no attrition among the 96 participants who attended the program. The 10 day program began on a Tuesday and finished on the Thursday of the following week (Table 1). Our 5 day, Monday to Friday, pilot programs proved too congested and caregivers requested that a weekend be included. This allowed caregivers time to spend talking together, having fun, such as a picnic, and consolidating some of the knowledge previously presented.

A major aim of the course was for participants to enjoy themselves. Sadly, fun and spontaneity are often lacking from caregivers' lives. Leisure pursuits, such as walks, table games like “Trivial Pursuit”, carpet bowls, singalongs, dances and going out for a drink, were included as part of the evening and weekend program. During these activities, caregivers would practise their

Table 1 Timetable of Dementia Caregivers' Training Program

	Morning	Afternoon	Evening
Day 1	Admission procedures Welcome and orientation	"Getting to know you" Reducing carer distress 1	Socializing
Day 2	Stress management and relaxation "Telling your story" Reducing carer distress 2	Healthy eating for older people	Film night
Day 3	Relaxation Re-roling	Reminiscence Keeping fit and healthy	Carer outing
Day 4	Relaxation Assertiveness training 1	Therapeutic use of activities and activity analysis Medical aspects of dementia 1	Socializing
Day 5	Relaxation practice with tape Picnic outing		Socializing
Day 6	Relaxation practice with tape Church	Sunday drive	Socializing
Day 7	Relaxation Communication	Assertiveness training 2 Medical aspects of dementia 2	Extended family sessions
Day 8	Stress management and relaxation Reality orientation	Work simplification and organization in the home Combating burnout	Extended family sessions
Day 9	Relaxation Use of community services	Coping with physical frailty	Socializing
Day 10	Relaxation "What if"—planning for the future	Farewell afternoon tea and presentation of diplomas	

skills of communication, activity analysis, reality orientation and reminiscence.

Patients and caregivers were given name tags and briefed daily after breakfast to review the previous night and to confirm each day's arrangements. Less threatening sessions were scheduled for the first week of the program; those that required more self-examination or were more confronting about the realities of the dementia were left until the second week. At the end of the program there was an afternoon tea graduation ceremony and presentation of a diploma.

Follow-through

Follow-through was an essential part of the program. While the 10 day program was both intensive and comprehensive and supplemented by take-home written materials and audiotapes, it was felt that the lessons would be lost without reinforcement. Telephone conferences were arranged with the coordinator at increasing time intervals over 12 months, starting at 2 weekly and finishing at 6 weekly intervals. The coordinator's input was gradually diminished on these telelinks. Towards the end of the year, the coordinator would absent herself from teleconference calls. Cohorts of caregivers attended the hospital for follow-up assessments at 3, 6 and 12 months after the completion of the program, thus providing opportunities for reunions. Relationships among cohorts of caregivers varied, with some establishing quite close friendships and continuing to meet informally at each other's houses. After the first year, annual telephone follow-ups were conducted. Two long-term outcomes were monitored, nursing home admission and death. Data on these endpoints were obtained for all patients.

THE TRIAL

We recruited subjects by seeking referrals from doctors, aged care and healthcare providers or by having articles in the

media. While the pilot programs had been heavily subscribed, once the research proper began, the well-known "disappearing subjects" phenomenon was evident. Of the 96 caregivers who entered the study, 40 did so of their own initiative, 16 were referred by local doctors, 15 through the Alzheimer's Disease and Related Disorders Society, eight after media publicity, and 17 through other sources.

To be eligible, we required that patients be less than 80 years old (for follow-up purposes) and have mild to moderate dementia, defined by the *Diagnostic and Statistical Manual of Mental Disorders*, 3rd edn¹¹. The patients had to live in a private home with a supporter, be able to understand English, not be a wanderer and not be aggressive. We stipulated that patients had to have dementia of mild to moderate severity defined as a score on the basic Activities of Daily Living of 0 or 1. All subjects agreed to random allocation to either a memory-retraining program (for patients only) or a dementia caregivers' program (as described above). Subjects were told that neither program offered a cure, but both offered the possibility of improvement in function. Institutional ethics committee approval was received prior to the trial and all subjects gave written informed consent.

Procedures

Prospective applicants for the program were sent questionnaires and the postal date on the envelope of the questionnaire determined which group eligible subjects were assigned to. Importantly, no systematic bias could have occurred because the time when caregivers sent their application was purely random and therefore meant that their allocation was not biased by any selection procedure. Subjects were assigned in turn to one of three treatment groups: *memory retraining program*, *immediate dementia caregivers' program*, or *wait-list or delayed dementia caregivers' program*. In the *memory retraining program*, patients were admitted for 10 days and received the patient component of the dementia caregivers' program, while

their caregivers had 10 days' respite. Those in the *delayed training group* received the dementia caregivers' program after approximately 6 months. We calculated a sample size of over 30 for each of the three groups as necessary for an intervention of moderate power (estimate 0.6) to produce a relevant effect size (estimate 0.67) for $\alpha = 0.5$.

Assessment

Patients were assessed on the Orientation Information Memory Concentration scale [OIMC, range 0–37 (37 = maximum cognitive functioning)¹²]; the Dementia Scale [range = 0–27 (higher score indicates worse function)¹²]; the Mini-Mental State Examination [range 0–30 (30 = maximum cognitive function, <17 indicated important deterioration)¹³]; the 21 item Problem Behaviour Checklist¹⁴ [range 0–42 (0 = no problems, 42 = all problems occurring frequently)]; the Activities of Daily Living [range 0–6 (0 = completely independent, 6 = completely dependent)¹⁵]; the Instrumental Activities of Daily Living [range 1–4 (1 = complete independence, higher scores indicate increasing dependence)¹⁶]; the 21 item Hamilton Rating Scale for Depression [range 0–64 (>16 indicates important depression)¹⁷]; the Geriatric Depression Scale [range 0–20 (>10 indicates possible depression)¹⁸]; and the Clinical Dementia Rating Scale for Dementia [range 0–3; (0 = healthy, 3 = severely demented)¹⁹].

Carers completed the General Health Questionnaire [range 0–30 (those with scores >4 probably were considered to have significantly psychological morbidity)²⁰]; and the Zung Depression Scale [range 20–80 (≥ 40 indicated important depression)²¹] and were rated on the Hamilton Depression Rating Scale. They were asked to keep a health diary of all the healthcare visits made and medications taken by them and the patient over the 12 months of follow-up. They were also asked to keep a record of all visits to day centres and any days in residential care. Completion of the diaries was encouraged at the regular telephone conference sessions. Demographic data on all participants were collected and included the position on the Congalton scale for socioeconomic status [range 1–7 (1 = highest status occupation, 7 = lowest)²²].

RESULTS

The Sample Defined

One hundred and one pairs of patients and caregivers entered the trial. Three of the 36 pairs in the *immediate carers' program* had insufficient follow-up data, and one of the 32 patients in the *memory retraining group* (MRP) changed caregivers during the follow-up period. These four pairs were excluded from further analyses. Two pairs in the *delayed carers' program* (DCP) completed intake and pretraining assessments but elected not to proceed with the training. Their data from the initial two assessments were included in the analyses at 0 and 6 months, but not subsequently. Otherwise all patients and caregivers who began the programs completed them.

This left 33 pairs in the immediate DCP, 32 in the wait-list DCP and 31 in the MRP. Of the 96 carers, 89 were spouses, four were siblings and three were children of patients. Forty-four caregivers were men and 30 of the 87 who completed the question affirmed their membership of the Alzheimer's Disease and Related Disorders Society. The caregivers' mean age was 67.7 (SD = 8.2 years).

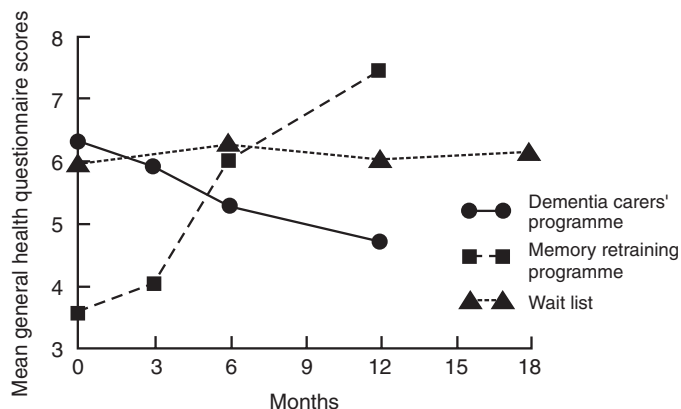


Figure 2 Mean General Health Questionnaire scores for carers in all three groups. Standard deviations of each group at zero, three, six, and twelve months respectively were 6.2, 5.8, 6.2, and 5.6 for the dementia carers' programme group; 6.3, 7.1, 8.4, and 9.4 for the memory retraining group; and 6.1, not available, 6.7, 6.6, and 7.7, at 18 months, for the wait list group. From Brodaty and Gresham², by permission of the BMJ Publishing Group

Of the 96 patients, 50 were men, 70 had probable Alzheimer's disease, 19 had multi-infarct dementia, and seven had other causes of dementia. Their average age was 70.2 (SD = 6.5 years; range 49–79 years) and they had had 10.4 (3.6) years' education and had mild-moderate dementia [Clinical Dementia Rating Scale score 1.1 (0.5)]. The duration of dementia, a mean of 3.8 (3.8) years, was similar in the three groups. The sample was predominantly middle-class ($n = 52$), with 16 from the upper socioeconomic classes and 25 from the lower socioeconomic classes. Data from three patients were missing. There were no significant differences between the three groups, for caregivers or patients, on any socioeconomic variable or initial measure of outcome at entry into the trial.

In later reports from the study, three subjects were excluded. One subject did not decline and he was subsequently rediagnosed as having benign forgetfulness; another subject who had undertaken the *memory retraining program* was excluded because he and his wife subsequently undertook a caregiver training program; and a third subject from the wait-list group did not provide sufficient data. Otherwise, all subjects declined over time, confirming their diagnosis of a progressive dementia. Diagnoses were able to be refined over time, so that of the 93 patients, 65 were subsequently diagnosed with probable Alzheimer's disease, 21 with multi-infarct dementia, three with Pick's disease and four with other uncertain cause of dementia (two subcortical dementia, one carbon monoxide poisoning and one diagnosis deferred). There were some slight differences in the baseline characteristics of patients and caregivers once these three pairs had been excluded, but these were trivial. Details can be found in the report from Brodaty *et al.*¹

Caregivers' Outcome

Caregivers' psychological morbidity, as judged by the General Health Questionnaire (GHQ-30) declined significantly over 12 months in the immediate intervention but rose steadily in the memory retraining group. GHQ scores of those in the delayed training program remained steady (Figure 2). Scores on the Zung Depression Scale did not show this differential effect, probably reflecting the low initial scores on that scale and the biological nature of many of its items²¹.

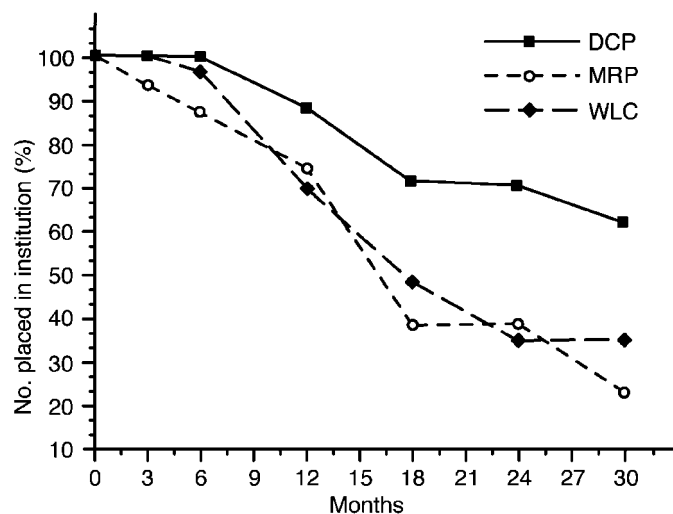


Figure 3 Survival curve showing percentage of patients not placed in an institution over time in the three programme groups. From Brodaty and Gresham², by permission of the BMJ Publishing Group

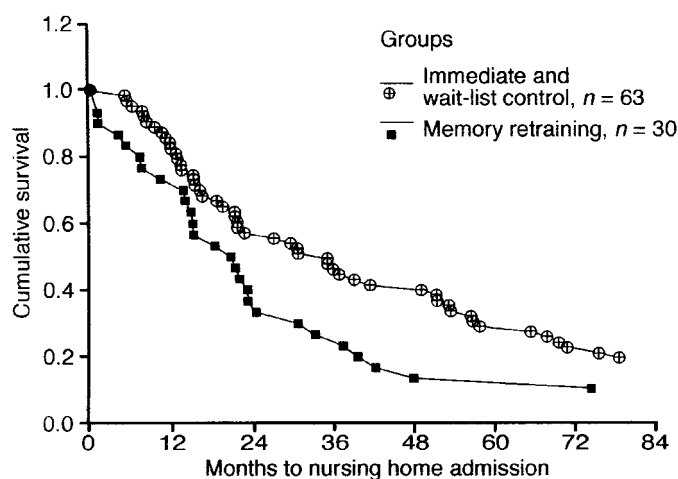


Figure 4 Kaplan-Meier survival functions for nursing home admission comparing the combined training groups with the memory retraining group. From Brodaty *et al.*¹, with permission. Copyright John Wiley & Sons Ltd

Patients' Outcomes

Over the 12 months there was a steady decline in all measures of patient cognition and function (Figure 2). Thus for the total sample, the MMSE scores declined from 17.1 (6.5) at baseline to 16.2 (7.3) at 6 months and 15.2 (7.6) at 12 months. Similarly, the Blessed dementia scale score increased from 7.0 (2.9) to 8.2 (4.3) and 10.4 (5.4) over the 12 months. Activities of Daily Living declined from 0.3 (0.6) to 1.1 (1.4) and 1.7 (1.7) over the 12 months. There were no differences between the three groups in the rate of patient decline. Patients were not depressed clinically. Fewer than six at any assessment over the 12 months had a Hamilton score of ≥ 16 . Their mean Hamilton and Geriatric Depression scale scores remained low and stable over time.

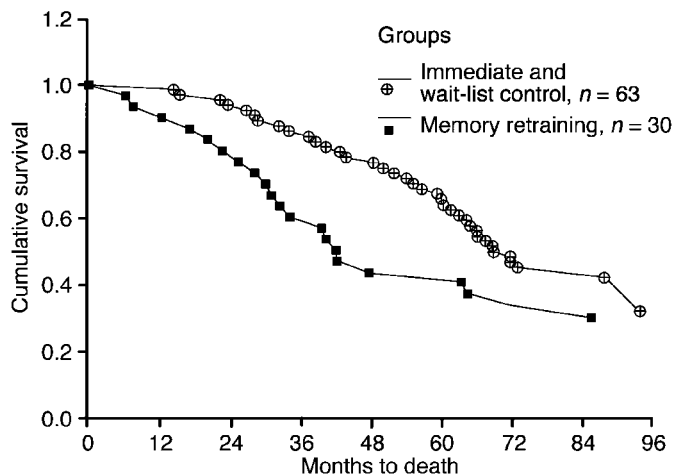


Figure 5 Kaplan-Meier survival functions for death comparing the combined training groups with the memory retraining group. From Brodaty *et al.*¹, with permission. Copyright John Wiley & Sons Ltd

Nursing Home Admission and Mortality

In the first 30 months, there was a marked difference between the groups in the percentage of patients still at home (Lee-Desu statistic = 6.19, $df = 2$, $p < 0.05$) (Figure 3). At 8 years follow-up, the rates of nursing home admission in the immediate and delayed carer training groups were similar and were combined, allowing for a comparison of the effect of caregiver training in general to the memory retraining group (Figure 4). Using Kaplan-Meier survival functions, there was a significant difference between the groups in survival at home and mortality. The rates of death of patients at 30 month and 8 year reviews were lower where caregivers had had training (Figure 5).

Costs

Program Costs and Institutional Costs

As the program was conducted within the psychiatric unit of a general teaching hospital, it was more expensive than may have been necessary. Even so, we were able to demonstrate that the costs of training were more than counterbalanced by the delay in nursing home admission. By 39 months follow-up, there were only 32 patients still alive at home: 17 of 31 (55%) from the immediate carers' program, 11 of 29 (38%) from the delayed carers' program and 4 of 30 (13%) from the memory retraining program. At the time, the average cost of a nursing home bed was *\$92.23/day. We calculated the total institutional costs for the three groups as \$19 918 for immediate carers' training patients, \$27 375 for delayed carers' program patients, and \$36 753 for memory retraining program patients. The cost of the training was estimated in 1991 as \$8868, including the hospital stay and 12 months' follow-up. This represented a saving of \$7967 (Australian) (\$5975 US) per couple in the immediate training program (compared to those in the memory training program) in the first 39 months of the program.

*Costs quoted in Australian dollars throughout. \$1 Australian = \$0.75 US at that time.

Health Care and Lost Employment Costs

We found that over the first 12 months of follow-up there were no appreciable differences between the groups in the number of visits to all doctors, general practitioners or non-medical health practitioners; or in the use of medication or of hospitals. In the second 6 months of the first year, memory retraining patients spent more nights in institutional care than the other two groups.

Prediction of Nursing Home Admission and Death

By 5 years follow-up, 75.8% of patients had entered a nursing home and 41.8% had died. Dementia severity and rate of deterioration and carer psychological morbidity significantly influenced rates of nursing home admission and death. These rates were comparable to previous reports^{23,24}.

Carer training had a significant protective effect against nursing home admission and, surprisingly (and independently of nursing home admission), against death. There was an association between earlier nursing home admission and caregiver distress, as measured by the GHQ; index measures of dementia severity and problem behaviours; more rapid decline in cognitive function; more rapid decline in overall dementia severity; and increase in problem behaviours.

COMMENTS

Carer intervention programs have considerable potential. They can improve the quality of life of the carers and probably that of patients¹. We have recently reviewed 35 controlled studies of carer intervention and found that a minority of them demonstrated clinically significant beneficial effects⁸.

Limitations to previous studies included: heterogeneity of patients and of carers in the sample; variety of recruitment methods; ceiling and floor effects as regards the outcome measures; low number of numbers and insufficient power; lack of blindness; insufficient duration of follow-up; and lack of specificity in matching interventions with carer needs⁸.

The Sydney Carer Training Program study overcame many of these limitations and demonstrated psychological improvement in carers, delay in nursing home admissions and cost savings. The delay in institutionalization was not at the expense of increased carer distress.

There were a number of unanswered questions from the study. It was not possible to know which components of the package of interventions were effective. We provided a broad-spectrum intervention—something for everybody. This was confirmed at exit interviews after 12 months' follow-up, where each component of the program was identified by at least one carer as being helpful to him/her.

The program was unnecessarily expensive in that it was conducted within a hospital setting. While this provided many advantages, it was very costly, and the cost analyses allowed for 20 hospital bed-days per patient-carer couple. We do not know whether the program needed to be residential, although this did provide some advantages. The advantages of residential programs are that they promote more cohesive bonding and allow for observation of behaviours not easily accessible within a day program. Clearly, residential programs could be conducted in less expensive settings.

The ideal number of couples per training cohort is unknown, but our impression was that numbers greater than 10 would impede the group process. Also, our experience suggested that the earlier the intervention, the

better, and that matching carer cohorts, e.g. spouses, younger people with dementia, socioeconomically and geographically, may have advantages.

Future research might benefit from a more targeted, selective approach—matching the needs of carers with appropriate interventions. The questions of which carer interventions benefit which carers for which patients at what time in the course of the dementia are complex. Finally, the advent of specific drug treatments for Alzheimer's disease begs the question of whether carer interventions plus drug treatments are superior to either alone.

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Elder Abuse— Epidemiology, Recognition and Management

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During the 1980s, elder abuse emerged as a health and social issue of international importance¹. It is defined as “a single or repeated act or lack of appropriate action occurring within any relationship where there is an expectation of trust which causes harm or distress to an older person”². While most authorities include self-neglect within the broad definition, acts which threaten an elder’s well-being as a consequence of their competently made decisions are specifically excluded.

Abuse may occur in one of two settings. Domestic abuse is perpetrated within the home of the victim or a caregiver by either a relative or other care provider. Institutional abuse occurs within a designated care facility (residential or nursing home or hospital), perpetrated by one or more individuals having an obligation to care for and protect the victim.

Five major categories of abuse have been identified^{3,4}:

1. *Physical*: any activity involving force to generate bodily injury or pain, including striking or burning and the use of physical or pharmacological restraint.
2. *Sexual*: any form of non-consensual sexual contact, including unwanted touching, rape, sodomy and coerced nudity.
3. *Psychological*: the infliction of distress through verbal or non-verbal acts, including insults, threats, humiliation, infantilization and harassment.
4. *Financial*: the improper use of an elder’s property or assets, including theft, deception, coercion and misuse of authority to act, such as power of attorney.
5. *Neglect*: the refusal or failure to fulfil care obligations, including the provision of food, water, clothing, medication, comfort and protection.

PREVALENCE AND INCIDENCE

Variability in case definition obscures direct comparison, although the prevalence of elder abuse is broadly similar throughout Europe and North America. To date there are few data from developing countries (Table 138c.1). With the exception of the USA, a lack of national incidence data reflects widespread absence of formal mechanisms for case reporting and validation. American data estimate the incidence of domestic abuse at 450 000 elderly people/year, of which only 16% are reported to statutory agencies⁵.

VICTIM CHARACTERISTICS

Likelihood of being abused increases with age⁶. Elders aged over 85 are at particular risk of neglect⁷, financial abuse⁸ and abuse by

designated carers⁹. There is some evidence that minority ethnicity and non-White race represent risk factors for abuse^{6,10}. Women are more likely than men to be victims¹¹ but this observation may be confounded by greater likelihood of living alone, which is associated with financial abuse⁸. In contrast, victims of physical, sexual or psychological abuse are likely to live with others, particularly a spouse or child¹². Poverty elevates the risk of abuse⁶.

The role of cognitive impairment is complex. Factors predicting abuse of dementia patients include behavioural disturbance, poor premorbid relationship with a carer and psychological or physical abuse by the patient¹³. The severity of cognitive impairment does not appear to be associated with abuse. Spouses caring for dementia patients are at particular risk of psychological and physical abuse³⁵. In one series, one-third of carers reported physical abuse by the patient, which in turn was associated with abuse of the patient by the carer¹⁴.

Vulnerability to abuse has been associated with certain personality traits¹⁵. Victims of psychological abuse have less ability to control problem situations and tend to react aggressively when feeling anger or frustration. In contrast, physical abuse victims pursue passive or avoidant behaviour, while financial abuse victims possess negative beliefs of self-efficacy and turn aggression or frustration on themselves.

ABUSER CHARACTERISTICS

Greater understanding of abusive situations has focused attention on those perpetrating abuse. Carers who suffer social isolation, feel unsupported and are financially dependent are at risk of abusing^{13,16,17}. Men are more likely to abuse than women, and more likely to cause physical abuse¹⁸. Psychological abuse is more likely to be caused by women. Abusers tend to suffer declining health and mental illness increases the risk of perpetrating abuse¹⁹. In particular, depression and anxiety among carers are associated with abusive behaviour^{20,21}.

The role of drug and alcohol misuse is controversial. Abusers identified as misusing substances are likely to be male children of the victim, less likely to provide care and more likely to cause physical or psychological abuse than financial abuse²². However, in one study of referrals to a community psychiatric service, consumption of alcohol by the carer was not associated with abuse¹³.

Table 138c.1 Prevalence (%) of elder abuse (aged 65+)

Abuse type	UK ¹	The Netherlands ²	USA ³	Canada ⁴	Sri Lanka ^{5*}
Physical	1.5	1.2	2.0	0.5	1.5
Psychological	5.4	3.2	1.1	1.4	1.5
Financial	1.5	1.4	–	2.5	1.0
Neglect	–	0.2	0.4	0.4	5.6

¹Ogg and Bennett³².²Comjis *et al.*³¹.³Pillemer and Finklehor¹².⁴Podnieks³³.⁵Lekamwasam and Chandanee³⁴.

*Aged 60+.

Table 138c.2 Risk factors for abuse

Victim
Age > 75
Living alone (financial abuse)
Living with spouse or child (physical, sexual or psychological abuse)
Low income
Cognitive impairment
Poor pre-morbid relationship with carer
Abusive, passive or avoidant behaviour
Abuser
Male (physical abuse)
Female (psychological abuse)
Social isolation
Financial dependency
Impaired physical or mental health
Substance abuse
History of receiving or perpetrating abuse

NATURAL HISTORY OF ABUSE

Those who have suffered abuse are themselves likely to become abusers²³ and carers who have violent elders express more violence to their dependants²⁴. In one follow-up study, 20% of victims suffered physical or financial damage following abuse, although 70% were able to stop abuse either themselves or with the help of others³¹. Corroborated abuse is associated with greater risk of death for elderly victims (odds ratio 3.1; 95% CI, 1.4–6.7) after adjusting for co-morbid and demographic factors²⁵. However, abuse rarely leads to homicide: in one study only 2% of elderly homicides could be attributed to abuse²⁶.

RECOGNITION

Diagnosis of elder abuse requires a high index of suspicion. Professionals should be alert to the presence of one or more risk factors for abuse (Table 138c.2) and sensitive to principal abuse signals (Table 138c.3). In cases of suspected abuse, a coordinated multi-agency approach must identify all care needs and deficiencies. Corroborated history must be obtained from all participants, including victim, alleged abuser and designated carers, with careful verification of information obtained from cognitively impaired individuals.

Symptoms and signs of abuse must be elicited during a comprehensive clinical assessment to which the victim consents (Table 138c.4). Accurate documentation, including note keeping, radiology and photography, will facilitate future management planning. The presence or absence of particular features do not alone confirm or exclude the diagnosis of abuse. Nevertheless,

Table 138c.3 Abuse signals²⁹

Caregiver personal problems
Caregiver interpersonal problems
Care receiver social support shortage and history of abuse

Table 138c.4 Symptoms and signs of abuse⁴

Physical abuse
Carer refusal to permit examination
Reports of being hit, kicked or mistreated
Unexplained behavioural disturbance
Presence of unexplained bruises, lacerations, ligature marks, fractures
Untreated injuries in various stages of healing
Inappropriate use of prescribed medication
Sexual abuse
Reports of sexual assault or rape
Bruising of breasts or genital area
Torn, stained or bloody underclothing
Unexplained genital infection or bleeding
Psychological abuse
Reports of verbal or emotional mistreatment
Withdrawal, non-communication or non-responsiveness
Unexplained or unusual agitation or behavioural disturbance
Financial abuse
Reports of financial exploitation
Unauthorized or unexplained changes in banking practice
Abrupt, unauthorized or unexplained changes to financial documentation
Unexplained disappearance of assets
Unmet care needs in the presence of adequate financial resources
Sudden appearance of individuals asserting their rights to an elder's assets
Neglect
Reports of mistreatment
Failure to provide food and hydration
Failure to meet clearly identified care needs
Hazardous or unsanitary living conditions

defensiveness and irritability by caregivers are predictive of abuse, poor physical care predicts physical abuse and psychosocial stress or exploitation predict psychological abuse²⁷.

MANAGEMENT

Denial, resistance to intervention, ignorance of intervention protocols, confidentiality and fear of reprisal have all been cited as professional barriers to the management of elder abuse²⁸. Central to effective management is the establishment of a single coordinating agency, providing education, advice and access to resources. Intervention models should be low-cost, multi-disciplinary, collaborative and capable of evaluating outcomes²⁹. The key elements of such a model may include:

- Educational resources.
- Mechanisms for accurate case identification.
- Professionals to identify and deal with health, social, financial and legal issues.
- Workers to support and monitor victims and their care networks.
- Victim empowerment and advocacy groups.

Identification of incipient abuse should generate a brisk response to avoid escalation. Management must aim to preserve

autonomous choice for the victim and avoid paternalistic action that seeks to provide a speedy resolution, perhaps through institutionalization. Accurate identification of unmet care needs should generate planned and effective care strategies, which engender safety without intrusion. Given that abuse may be multidirectional, attention may need to be focused on both victim and abuser.

While overtly criminal activity, such as theft or assault, may be dealt with by prevailing criminal law, most countries have eschewed a legislative approach to elder abuse, relying instead upon health and social service agencies to develop locally applicable policies and procedures. In the USA mandatory reporting laws have achieved only limited success and professionals remain unfamiliar with reporting procedures³⁰. Of greater potential benefit for the future is the emergence of national organizations, aiming to prevent the abuse of older people by disseminating research, informing public policy and providing specialist advice to both professionals and the general public³.

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The Care of the Dying Patient

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The care of the older dying patient is provided in many settings, including homes, tertiary care hospitals, nursing care facilities and inpatient hospices. The issues relevant to psychiatry are relatively consistent across these contexts. This chapter provides an overview of the major psychiatric syndromes and therapies that are covered more thoroughly in the primary palliative care literature.

PSYCHIATRIC SYNDROMES IN PALLIATIVE CARE

Depression

Depression is common in terminal illness. The prevalence of a major depressive syndrome in late-stage cancer patients has been estimated at 23–58%, varying by diagnostic criteria and methodology¹. The diagnosis of depression is sometimes difficult to make in the setting of medical illness, because many of the symptoms of the patient's condition may overlap with those of depression, such as fatigue, loss of energy and altered sleep and appetite. Furthermore, many of these same symptoms can be side effects of medications, such as corticosteroids or chemotherapeutic agents. The general approach to this diagnostic problem is to rely more on the psychological and cognitive symptoms of depression, such as worthlessness, hopelessness, excessive guilt and suicidal ideation². One method used to clarify the distinction between depression and medical illness is the substitution of physical symptoms of uncertain etiology with psychological counterparts. An example of this approach is found in the Endicott Substitution Criteria³, which replaces physical symptoms as follows: (a) change in appetite and weight is replaced by tearfulness and depressed appearance; (b) sleep disturbance is exchanged for social withdrawal and decreased talkativeness; (c) fatigue and loss of energy are substituted with brooding, self-pity and pessimism; and (d) diminished concentration and indecisiveness are replaced by lack of reactivity.

Another complexity in diagnosing depression in dying older adults is the question of how much of a patient's symptoms should be attributed to the normal difficulties of adjusting to the terminal nature of his/her diagnosis. When interviewing the patient, each of the clinical symptoms of depression should be explored in detail, with special attention to its pervasiveness and its interference in functioning. For example, hopelessness may be understandable with regards to hoping for a cure or recovery, but patients who are not depressed may be able to maintain hope for other improvements, such as better symptom control, or a better connection with their loved ones before their death. Hopelessness that is accompanied by despair or suicidal ideation is highly associated with a depressive syndrome and usually merits

treatment⁴. While the multitude of issues relevant to its discussion is beyond the scope of this chapter, the assisted suicide movement heightens the need for careful assessment of depressive symptoms in the elderly dying patient.

The treatment of depression in terminally ill older patients often includes pharmacotherapy and is similar to other geriatric settings in its attention to managing symptoms and side effects. For this reason, selective serotonin reuptake inhibitors (SSRIs) are widely used in this setting because of their tolerability and their ability to also address anxiety. However, the principal side effect of SSRIs, gastrointestinal upset, can lead to weight loss, which is particularly detrimental to debilitated, medically fragile patients. Special care needs to be taken regarding drug interactions of the cytochrome P450 system with this class of drugs. Venlafaxine, a serotonin–norepinephrine reuptake inhibitor (SNRI), is less likely to interact with other medications because of its low protein binding (35%). Terminal illness makes a second variable relevant in medication choice, that of likely duration of treatment. In the setting of a life expectancy of days to weeks, an SSRI will not have time to take effect. For this reason, many clinicians favor the shorter-acting SSRIs, such as sertraline or citalopram. Bupropion has a similarly low side-effect profile as the SSRIs and also has some of the same dopamine agonist properties as the psychostimulants (see below), which may make it a good alternative in this setting. Tricyclic antidepressants are useful as adjuvant pain control and for sleep, but their side-effect profile and drug interactions make them a second-line choice.

Psychostimulants are a well-studied alternative for treating depression in terminally ill patients. While they are less useful in general populations because of tolerance and abuse potential, patients at the end of life can tolerate stimulants for up to a year without significant abuse problems². Particularly helpful in the depressed patient with psychomotor retardation and mild cognitive impairment, stimulants such as methylphenidate and dextroamphetamine have been shown to reduce sedation from narcotic analgesics and also have adjuvant analgesic effects in cancer patients⁵. Pemoline is a less potent stimulant with a lower abuse potential than methylphenidate, and therefore has the advantage of being less regulated by the federal government. Also, it is available in a chewable form that can be advantageous for patients who have trouble swallowing or have intestinal obstruction⁶.

Anxiety

As with depression, anxiety must be viewed as part of a normal spectrum in the patient's reaction to a life-shortening illness. Death can be seen as a model for human feelings of abandonment

and separation⁷ and therefore brings with it the likelihood of anxiety that needs to be addressed. Such anxiety can often interfere with interpersonal relationships and impair the patient's ability to understand and make decisions about his/her treatment. The range of etiologies for anxiety in the terminally ill is wide, and includes the full range of psychiatric disorders (adjustment disorder, panic, generalized anxiety disorder, phobia) but also covers a number of medical conditions such as hypoxia, sepsis, pain, akathisia, and withdrawal from alcohol, barbiturates or benzodiazepines⁸. One must also consider the restlessness and agitation that can be part of delirium. Each patient's anxiety must be assessed with empathy and a special care toward building rapport with the patient, since open discussion in itself can be very therapeutic in the patient who fears abandonment at the end of life.

When a patient's level of distress is sufficiently intense, treatment is warranted. In mild cases, cognitive-behavioral techniques, such as progressive muscle relaxation, can be very effective⁹. With more serious anxiety in terminally ill patients, pharmacotherapy is indicated. Benzodiazepines are a mainstay of anxiolytic treatment in the population, especially because of the decreased concern regarding abuse potential. Short-acting agents, such as lorazepam and alprazolam, are favored for older adults over their longer-acting counterparts, such as diazepam and clonazepam, because of the potential for the impaired metabolism in many elderly individuals to lead to toxic accumulation of drugs or their metabolites. Several of these medications are available parenterally, as is a rectal form of diazepam, which is widely used in the hospice setting. Neuroleptics, such as haloperidol and risperidone, tend to be favored when the etiology of anxiety is suspected to be due to corticosteroids or delirium, or if there are psychotic symptoms associated with the anxiety². Hydroxyzine is an antihistamine which is an effective sedative, anxiolytic and also has some analgesic properties and has minimal side effects^{10,11}. Lastly, opiates are very effective in reducing anxiety, especially when it is associated with pain. Their use, however, is limited by respiratory depression.

Delirium

Delirium is a disorder of arousal and cognition caused by a medical condition that has relatively rapid onset and fluctuating course. Delirium has a prevalence of one-quarter to one-half of patients at the end of life¹². This condition greatly complicates the treatment of the patient's other symptoms, such as pain, and causes great concern in family members and staff. In many cases delirium is reversible, even in relatively advanced terminal illness. Therefore, an investigation into reversible causes of delirium should be considered, including complete blood count, electrolyte studies, including calcium level and coagulation panel to rule out sepsis, dehydration, hypercalcemia and disseminated intravascular coagulation.

The diagnostic work-up of delirium in the terminally ill differs from that of the general population, in that special consideration must be made regarding the patient's or the family's wishes for invasive procedures to diagnose or treat the underlying cause of the delirium. The cause of terminal delirium most often is multifactorial or may not be found. Furthermore, when a distinct cause is found, it may be irreversible (such as hepatic failure or brain metastasis)². In one study, an etiology was discovered in less than 50% of terminally ill patients with delirium¹³. Lastly, it should be noted that in the last 24–48 h before death, cognitive changes are usually irreversible and are attributable to multiple organ failure.

Treatment of delirium involves correction of any reversible underlying causes that can be found; supportive therapies, such as fluid, nutrition and vitamins are often helpful. Non-pharmacolo-

gic interventions to help reduce anxiety and disorientation include a quiet, well-lit room with familiar objects, a visible clock or calendar and the presence of family¹². When these measures are insufficient, neuroleptics are the treatment of choice, especially in patients with an agitated presentation. Haloperidol in low doses is still favored by many clinicians; intravenous administration can hasten the onset of action, and may be less apt to cause extrapyramidal side effects than the oral form. If sedation is necessary, a short-acting benzodiazepine can be added.

PSYCHOSOCIAL INTERVENTIONS IN PALLIATIVE CARE

Even in the absence of serious psychiatric morbidity, the end of life can be a period of considerable intrapsychic and interpersonal conflict for the patient. In many cases, psychotherapeutic interventions can offer considerable benefits to a palliative care setting. The fact that this area of intervention is not better studied may reflect anxieties in therapists who are forced to confront their own mortality in providing palliative care¹⁴. Methods vary in the treatment of dying patients, but some common elements unite the different approaches. In general, psychotherapy with patients with terminal illness should seek to:

1. *Allow the patient to openly communicate his/her feelings about death.* The dying patient experiences many fears, and often feels isolated because there is no-one with whom they can be shared¹⁵.
2. *Strengthen the interpersonal bonds with those who will survive the patient.* This serves to alleviate the sense of abandonment that accompanies dying and allows the patient a sense of having "put affairs in order". It is also important for the survivors and their eventual grieving process, since ambivalent feelings toward the deceased are a risk factor for complicated bereavement¹⁶.
3. *Establish hope while gently confronting denial.* While hope of cure may not be realistic, it can be replaced by the hope to live to be present at certain near-term events such as a graduation. Similarly, patients can hope for dignity in their final days, for freedom from pain or for their personal understanding of an afterlife.
4. *Help the patient find existential meaning and coherence in his life and in his death.*

Kubler-Ross

Any discussion of the psychology of dying owes a debt to Ellen Kubler-Ross and her seminal book, *On Death and Dying*¹⁷. Kubler-Ross interviewed hundreds of patients with terminal illness and described the process of dying in five stages: (a) *denial and isolation*, in which the patient refuses to believe he/she is terminally ill and feels suddenly alienated from the rest of the population that is not acutely dying; (b) *anger*, which can be expressed either overtly or in subtle, passive ways, depending on the personality of the individual; (c) *bargaining*, in which the patient seeks to prolong life by making promises to a higher power; (d) *depression*, the sense of existential loss upon realizing that death is unavoidable; (e) *acceptance*, in which the patient achieves a serenity and grace even in the face of death and comes to terms with the inevitable. Kubler-Ross herself states that these stages may recur several times and vary in order throughout the patient's terminal illness, and any one stage may be missing altogether. Therefore, these stages should only serve as guideposts to anchor one's clinical assessment and interpretation of the dying

patient. Any attempt by the therapist to impose the stages as a structure can be an undue burden placed on the dying.

Psychotherapeutic Care

Psychotherapeutic approaches vary widely but, in general, the psychodynamic modes employ open-ended methods through which the patient gains insight. Underlying this approach is the belief that by better understanding the emotional pain of terminal illness, the patient will gain some relief from it¹⁵. The therapist pays careful attention to the patient's defenses, such as denial, displacement, counterdependency and dependency, and those that are felt to be adaptive are gently reinforced. If the defense is maladaptive, the question becomes whether the patient could tolerate an attempt to change it. Denial is felt to be the most common defense encountered in clinical practice, and it can be the most difficult to navigate for those who do not commonly work with the dying. The decision of whether or not to confront denial plagues psychiatrists, primary physicians and families alike. Stedeford¹⁵ recommends that denial is a problem only when it is the sole or prominent defense. It then blocks communication with family and friends and prevents making suitable plans for the future. Often denial serves an important role in allowing the patient time to assimilate gradually information that would be overwhelming if absorbed all at once. Connor¹⁸ believes that most denial is used to preserve interpersonal relationships. In this model, patients use denial primarily to cope with guilt about the effect of their condition on others, to protect others from the emotional stress they might feel if the patient were to openly acknowledge his/her condition and feelings, or out of fear of abandonment. He devised an intervention for this type of denial through a structured set of questions displayed in Table 138d.1.

Some other techniques are especially appropriate with dying older patients. Problem-solving therapy is a brief treatment that attempts to alleviate emotional symptoms by focusing on the social and practical difficulties faced by patients¹⁹. These problems are linked to the patient's symptoms, and the patient is helped to solve the problem by breaking it down into stages as follows: (a) clarification and definition of the problem; (b) setting of achievable goals; (c) consideration of alternative solutions; (d) selection of a preferred solution; (e) clarification of the necessary steps to implement the solution; (f) evaluation of progress. The advantages of this therapy are its brief format, which is sometimes necessary to fit the time frame of dying patients, as well as its accessibility to patients, who are sometimes

uncomfortable with the jargon of psychotherapy. Cognitive and behavioral techniques are widely used to address specific symptoms, such as anxiety and phobias, through progressive muscle relaxation, imagery exercises or cognitive restructuring⁹. Guided imagery and trance states have also been successfully used to treat cancer pain. Education is often overlooked as a highly therapeutic tool to combat anxiety, since much of a dying patient's fear is generated by the unknown. The resources to assist in this education of the patient and family are growing exponentially and are available through varied media.

Psychological Benefits of Hospice Care

The first modern hospice was founded in 1967 in England to address concerns about the poor training of physicians to deal with terminal illness. Since then, the hospice movement has grown and spread to the USA, where roughly 20% of all deaths are now accounted for by patients who use hospices. The hospice approach seeks to improve the quality of the end of life by focusing on the whole patient, including his/her medical, pain relief, emotional and spiritual needs. Hospices employ a number of the above described therapeutic interventions through varied disciplines with a distinctive commitment to a team-based approach. Along with the psychological benefits for the dying patient in receiving this interdisciplinary, psychologically sensitive care at the end of life, there appears to be a psychological benefit to survivors of patients who use hospices; McNeilly and Hillary found survivors of hospice patients less likely to regret not having more openly expressed feelings to the person they cared for than those who used a home health care group²⁰. Hospices can be seen as an intentional aid to the patient in his/her psychological journey at the end of life in its explicit commitment to helping the patient "die well", as he/she understands and interprets it, while maintaining an abiding commitment to the family and other survivors.

Although psychiatric care of the dying older patient embodies many challenges, it is full of rewards for the physician, patient and the patient's families. The reciprocity inherent within the relationship with dying patients is profound in its implications for educating us about life, suffering and adaptation. The more capable we become of caring well for the dying during this transition in their lives, the better we will be at understanding and caring for all of our patients.

Table 138d.1 Structured psychosocial intervention for denial

1. Different people experience different kinds of difficulties when they are ill. What, for you, have been some of the most difficult aspects of having your illness?
2. Are there any things you do, or that other people do, that make these difficulties easier to deal with?
3. Is there anything you or other people do, that make these difficulties harder to deal with?
4. Do you believe you will or will not recover from this illness?
5. Have you had any close encounters with death?
6. What effect has your illness had on your family and close friends, and how have they reacted to it?
7. How do you feel about the way in which your family and friends have been affected by or have reacted to your illness?
8. Is there anything good that has come out of your having your illness?
9. Are there any other thoughts or feelings about your illness or the questions I've asked that you'd like to talk more about?

From Connor¹⁸, with permission.

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Prevention in Mental Disorders of Late Life

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INTRODUCTION

In its influential 1994 report, *Reducing Risks for Mental Disorders: Frontiers for Preventive Intervention Research*, the Institute of Medicine (IoM) of the US National Academy of Sciences¹ assessed the state of knowledge in prevention research and identified directions for future scientific development. The index of this substantial volume contains no entries for “aging”, “aged”, “elderly”, “geriatric” or “gerontological”. In a section on illustrative preventive intervention research programs, examples are given of two service programs, one for caregivers of patients with Alzheimer’s disease and one for widows, that are considered to have relevance to prevention. The authors could identify no randomized controlled preventive trials in the area of aging.

Prevention in the mental health field has been seen, traditionally, as an area that has been implicitly restricted to those concerns designed for application to issues in childhood and adolescence. If anything, prevention in geriatrics was seen as an oxymoron. Very simply, prevention was taken to mean youth. Theory and research in prevention were restricted to issues of child development and intervention early in the life course.

Why be concerned with prevention in late life? As is well covered in other sections of this text, there is the demographic imperative brought about by the overall aging of the world population and, in particular, by the aging of the older population. As pointed out in the classic papers by Gruenberg² and Kramer³, the same dynamics—public health measures, technological development and lifestyle changes—that created this growth in the overall population were also relevant to growth of the population of those with chronic illnesses and disabilities. They conclude that, in the absence of cures or effective preventive strategies, we will see an explosion in the number of older persons with serious and persistent disabling illnesses, particularly mental disorders. The availability of more efficacious treatments and the accessibility of appropriate services in the community combined to produce huge gains in the life expectancy of those with mental disorders who, in earlier times, would have died long before reaching old age. This demographic imperative leads to the conclusion that prevention must be an important part of the agenda of geriatric psychiatry.

The traditional public health view derives from infectious disease and is divided into primary, secondary and tertiary prevention. Primary prevention is directed toward maintaining health by isolating the causes of disease and eliminating or counteracting them. Secondary prevention is directed toward enhancing recovery by case identification and prompt intervention early in the course of illness. Tertiary prevention is directed toward those already ill and emphasizes treatment and rehabilitation⁴.

There is a growing consensus that the traditional public health view is not optimal. The components of this approach, including, for example, concepts such as pathogens, risk factors, disease vectors and definitions of caseness, do not translate easily into psychopathology or chronic disease. The 1994 IoM report¹ adapts a scheme developed by Gordon⁵ to characterize preventive interventions as universal (targeted to a general population), selective (targeted at individuals at increased risk) or indicated (targeted to individuals with minimal levels of signs or symptoms).

Universal interventions are broad public health measures intended for an entire population or for significant geographic, socioeconomic or categorical subgroups within it (e.g. rural residents, low-income older persons, or pregnant women). Universal interventions (e.g. iodizing salt) may reduce risk for a large segment of a population but in all likelihood do not have impact on those already at high risk or those who would not have been at risk at all. Cost–benefit assessment is a clear decision criterion for the development and implementation of universal interventions, since they would, by necessity, involve exposure of many individuals not at risk for development of an illness.

Selective interventions are targeted toward those individuals at significantly increased risk of developing the particular illness or condition. Genetic loading and positive family history, other illnesses, or psychosocial or environmental transitions are all examples of factors that might be used to target selectively preventive interventions. Although selective interventions may seem easy, careful efforts are needed to assure proper identification and to avoid stigmatization or unnecessary fear.

Indicated interventions are targeted on those individuals who are already symptomatic and in whom early intervention may alter the longitudinal course or optimize the outcome of the illness. The underlying assumption is that significant advantage is gained when preventive interventions are extended to concerns with function and disability in those who already have an illness. Interventions directed at minimizing post-treatment relapse or recurrence would also fit within this category⁴¹.

Following recommendations of the National Advisory Mental Health Council⁶, this chapter is based on assumptions that an appropriate approach to prevention must: (a) be tied closely to treatment; (b) have strong connections to service systems and services; and (c) be based upon models of etiology, pathophysiology and risk. Following Kraemer *et al.*⁷ we use “risk” and “risk factor” narrowly to indicate an empirically demonstrated agent or exposure that influences the likelihood of an event in a defined population. Preventive interventions are those directed at reducing risk of the development, exacerbation or adverse consequences of mental disorders.

PREVENTIVE INTERVENTIONS IN THE CONTEXT OF TREATMENT

Prevention of Relapse and Recurrence

Establishment of the efficacy of treatments is one of the major accomplishments of geriatric psychiatry research. As outlined in other sections of this text, many studies have demonstrated that older patients respond robustly to treatments that are appropriately applied with adequate intensities. These data are, largely, based on relatively brief, randomized controlled trials addressing the short-term efficacy of treatments to manage the symptoms of serious illnesses like depression⁸ and Alzheimer's disease^{9,10}. More recently, the prevention of relapse and recurrence has emerged as a major orientation in the treatment of the older patient. As the recognition has grown that most mental disorders in late life are chronic, recurring illnesses with substantial residual disability¹¹, so too has the acknowledgement that treatment must be approached with a much longer-term perspective¹². An intervention is preventive if acute treatment response is the starting point, with the major purpose of preventing relapse or recurrence and not managing symptoms.

Prevention of Side Effects and Adverse Reactions

Co-morbidity and the associated polypharmacy that comes from treatment of multiple conditions are characteristic of older patients. New information on the genetic basis of drug metabolism and on the action of drug-metabolizing enzymes now provide us with important perspectives on clinically significant alterations in drug concentration levels or on complex drug interactions¹³. For example, many of the newer antidepressant agents, the selective serotonin reuptake inhibitors (SSRIs), compete for the same metabolic pathway used by β -blockers, type I C antiarrhythmics and benzodiazepines.

Many older patients require antipsychotic treatment for the management of behavioral disturbance in schizophrenia, depression and Alzheimer's disease. Movement disorders are common side effects of the older types of these medications, the conventional neuroleptics. Although doses tend to be quite low compared to doses used in young or mid-life adults, age and length of treatment represent major risk factors for the development of movement disorders¹⁴. Recent data suggest the possibility that the newer antipsychotics present a much lower risk and that the development of tardive dyskinesia may be preventable through use of different medication¹⁵. Trials of agents (e.g. antioxidants) hypothesized to treat these side effects have been proposed, although the data from some of the early studies have been inconsistent^{16,17}.

Many drugs affect body sway and postural stability, although there is substantial variability within classes of drugs¹⁸. In older patients, where the prevention of falling is a major concern, a preventive strategy would reflect a differential selection of treatments or development of fall-specific preventive interventions¹⁹.

PREVENTIVE INTERVENTIONS IN THE CONTEXT OF SERVICES

Prevention of Suicide

Recognition of mental illness in older patients is highly variable. Changes in cognition, affect, thinking, sleep, etc. are often attributed to normal processes of age-related change by older people themselves, their spouses and close family members, and

even by their family doctors and primary care physicians. The most tragic result of the failure to recognize illness is suicide. Suicide rates increase with age, and men always outnumber women in suicide completion. In most countries, older men generally are at the greatest risk of suicide. In the USA, for example, old White men have a rate of suicide six times that of the general population. Psychological autopsy studies show that depression is common among these men but that it is rarely recognized. Nearly 40% of the men who kill themselves see their primary care doctors in the week of their death; nearly 70% in the month of their death²⁰. An uncontrolled field experiment on the island of Gotland in Sweden suggested that a depression-orientated educational intervention directed toward primary care physicians could reduce suicide²¹. Other approaches to the prevention of suicide, using aggressive outreach and case-finding techniques, have been developed in the context of community-based mental health or aging services^{22,23}.

Prevention of Premature Institutionalization

Nursing home placement typically comes at the end of a long and difficult period of caregiving by the families of patients with Alzheimer's disease. The burden of this caregiving in terms of stress, depression and quality of life has been extensively documented²⁴. It is only the rare (and very wealthy) family that can provide the care necessary to maintain a patient with Alzheimer's disease at home; institutional care is required for virtually all patients who survive to the end-stage of the disease. From a public health perspective, delay of institutional placement until it was absolutely necessary could have significant impact. In an important randomized controlled trial of a family-based counseling intervention, Mittelman *et al.*²⁵ demonstrated clear benefit: a delay of over 300 days in nursing home admission for patients whose families were randomized to receive the treatment. The counseling intervention also resulted in a significant reduction in depressive symptoms in these caregivers. Clinical drug trials in Alzheimer's disease have begun using institutionalization as a primary outcome²⁶ or as an outcome from open-label follow-up after the trial had ended²⁷.

Prevention of Excess Disability

The concept of excess disability is a classic one in geriatrics²⁸ and refers at its core to the observation that many older patients, particularly those with Alzheimer's disease, are more functionally impaired than would be expected on the basis of the stage or severity of their mental disorder. There are many sources of this excess disability: some are medical, some are psychosocial and some are environmental. A generation of research has clearly demonstrated that attention to these issues, and aggressive intervention where appropriate, will prevent excess disability and will optimize levels of function.

MODELS OF ETIOLOGY, PATHOPHYSIOLOGY AND RISK

Biological Models

Improved understanding of the etiology and pathophysiology of mental disorders can potentially lead to interventions that will prevent the onset or progression of disease. A useful model here is the large simple trial in a broad population; incident cases represent the primary outcome. The state of our knowledge is not yet sufficiently well developed to support this type of research.

Nonetheless, there are some interesting possibilities developing as we learn more about oxidative and inflammatory processes, apoptotic mechanisms, hormonal correlates and genetic factors in disease. One possible example is the area of vascular depression. Several different lines of evidence are now supporting the conclusion that one form of late-onset depression has a cerebrovascular etiology²⁹. Trials of vascular agents could use incident depression as outcomes.

Genetic Models

The genetics of mental disorders is an area of expanded activity³⁰. Notably, several genes are now implicated in different forms of Alzheimer's disease. At present, the genetic correlates of mental disorders are not sufficiently specific to be used for purposes of population screening. It is entirely conceivable, however, that trials directed at delaying onset of disease, or directed at minimizing excess disability, could be launched using some of this genetic information as a basis for subject selection.

Clinical Models

Co-morbidity is one of the hallmarks of mental disorders throughout the life course. In late-life mental disorders, the most frequently observed co-morbidities are physical illness and brain disease. This provides a possible opportunity for preventive interventions; we present a few examples.

Visual impairment is common among older people; there is a high frequency of depression among older persons with impaired vision, and the depression is more strongly predictive of disability than is the vision loss³¹. The vision clinic would seem to be an appropriate location for the development of programs orientated to the prevention of depression in older patients. Similarly, aggressive intervention in depression has the potential for minimizing disability and optimizing function. This latter point is supported by a broad range of studies demonstrating that depression compromises the outcome of rehabilitation in stroke, Parkinson's disease, heart disease, pulmonary disease and fractures³². Research has also demonstrated that treatment of depression can significantly improve outcome of treatment and rehabilitation for the co-morbid physical illness or condition^{33,34}.

Complex programmatic interventions have been shown to have efficacy in the area of prevention as well. Ray *et al.*¹⁹, for example, show how an intervention directed at safety with appropriate use of wheelchairs, psychotropic drugs and provision for transfer and ambulation, has a significant impact on the reduction of falls in nursing homes. In the outpatient setting, multidisciplinary comprehensive assessment resulted in significant improvements in the detection of depression and cognitive impairment and resulted in substantially reduced risk for nursing home placement^{35,36}.

Psychosocial Models

Bereavement is perhaps the prototypical psychosocial risk factor in late-life mental disorders³⁷. Considerable research has been devoted to the exploration of bereavement-related depression, complicated grief, traumatic grief and similar constructs³⁸⁻⁴⁰. It is entirely conceivable that preventive interventions for those with complicated grief could have major impact on both mental and physical health. It is also conceivable that greater understanding of the correlates and predictors of complicated or traumatic grief may lead to the design of earlier-stage interventions, directed at modifying the likelihood of developing the risk factor in the first

place. Interventions directed toward risk factors, rather than toward disorders, may be properly considered to be primary prevention.

CONCLUSION

In this chapter we have presented the broad parameters of a programmatic approach to prevention in late-life mental disorders. There is no established approach to prevention research that could be easily adapted to meet the needs of the field. Approaches to prevention need to be based in the deep foundation of knowledge in treatment and services research. They must be built upon improved models of etiology, pathophysiology and risk. A new approach to prevention holds the promise to be a significant development for the field. We have no doubt that success is achievable.

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A Damning Analysis of the Law and the Elderly Incompetent Patient—Rights, What Rights?

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Does the legal process provide a framework in the UK for protecting the rights of those who lack the capacity to make their own decisions? In what way should the law impact on families, professionals and carers? Surely you, the practitioner, you the carer, you the family member take decisions on what you believe to be in that person's best interests? You are there to do a good job exercising sound judgements. You are motivated by a desire to protect, to support and to make safe decisions in the best interests of the incapacitated person. You surely do not want to be impeded by considerations of an incapacitated person's legal rights. But what if the person him/herself wishes to make a different decision? You may consider it unwise or even risky for him/her to follow his/her desires. On what legal basis does one citizen override the ability of another to exercise choice? Who decides?

DOES THE LAW HAVE A ROLE?

Why is the law of incapacity/incompetence so important when it comes to protecting rights? For a person with capacity, the law recognizes the right to take responsibility for decisions. However, we must then bear the consequences of those decisions. Incapacity equates with society's view that incompetent adults may have to be protected. They may lack the ability to realize the consequences of the exercise of their decisions and that could place them at risk.

The dilemma is therefore apparent. At some point we cross that invisible divide between capacity and incapacity; between the right to make unsafe decisions and the right to be protected from them; between taking responsibility and the loss of the power to exercise our own judgement; between having enforceable rights and not.

For those who are deemed to lack capacity, the responsibility for deciding what is in their "best interests" may lie with those for whom there is a conflict of interest. They may be going to inherit the estate or they may even be those who are perpetrating abuse. The state of UK law leaves the incapacitated ripe for exploitation. Proposals for a change in the law appear to have stalled.

Perhaps when it comes to making a will, preparing an enduring Power of Attorney or protecting someone's financial affairs from fraud or exploitation, it is easy for us to accept the need for a protective framework. However, what about the concept of

enforceable human rights? When should the elderly be allowed to take risks?

We, ourselves, have rights and we can take steps to enforce them. But if we have lost the capacity to fight for or defend our rights, what then? When we have lost capacity, is there a right that restrains the distribution of our confidential information? "I don't want to go into a nursing home." "I want to choose with whom I live and with whom I have a personal/sexual relationship." "No, whatever they say, I do not believe that my children are acting in my best interests." Who will enforce my rights now? Who will listen?

On 2 October 2000, the UK Government belatedly brought into force the Human Rights Act 1998 in England. It sets out the rights and responsibilities of the citizen and provides a framework based on the European Convention of Human Rights and Fundamental Freedoms. It provides a baseline of rights below which the citizen should not be allowed to fall and introduces a rights culture laid on our domestic legislation. The indications are that it could have far-reaching effects on the practice of old age psychiatry, as well as impacting on the causes of action open to an increasingly aged population when their rights and opportunities are infringed.

The greatest impediment to the development of a rights-based system is the identification of the individual who will act on behalf of the incompetent client. Who will make the bridgehead to the lawyer? Who will blow the whistle? How does the incompetent client get to know of his/her rights in the first place? What happens when we cannot trust the families? The questions are easier to identify than the answers.

ROLE OF THE LAW

The role of the law must be viewed within the context within which it prevails. At present there is no doubt that in the UK^a the percentage of the population reaching the traditional retirement age^b is increasing and has done so since the post-war boom of the 1950s and 1960s. This, of course, has both political and financial ramifications. Linked to this has been a gradual increase in income, disposable assets such as homes, cars and other property, as well as a notable increase in the number of elderly people with their own assets.

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^aUnlike the Republic of Ireland, where the average age of the population is 28 years of age.

^b65 for men and 60 for women.

In examining a rights-based culture, it is relevant to remind ourselves that a notable shift emerged during the Conservative/Thatcherite years from 1979 onwards, when notions of solidarity and community waned and a “self” culture emerged.

What does all this mean in the context of the law, elderly people and incapacity? What does it mean within a society focused on “self” and where there are a greater number of assets and a greater number of people who need to have their rights protected?

Capacity is a concept based in our common law. Therefore, those with the final say are the judiciary. However, rarely is the law asked to be the arbiter of individuals’ capacity. Where there is dispute, it is always possible to seek a declaration from the courts. In practice, however, the final decision is taken by the myriad of individuals who come into contact with the incapacitated person. It could be the doctor, the nurse, the social worker, the staff in the nursing home or even the family. When it comes to medical treatment, the doctor who is in charge of treatment has the responsibility to make that decision. Some believe that only psychiatrists can perform this task. Psychiatrists get called in unnecessarily by other doctors (e.g. general practitioners, surgeons or anaesthetists) who do not seem to realize that they themselves are responsible. Usually those who are deciding on capacity have no concept of the test to be applied and merely assert that the individual will now have decisions made for him/her. It is a chilling scenario, particularly as many of us may eventually be in this predicament.

It is worrying that at times, medical staff, without any consideration of capacity, seek to obtain consent to a medical intervention from a person who does not have the capacity to consent. If that patient declines to agree with the treatment, then it is not given. It could perhaps have been justified by the common law doctrine of necessity. Failure to treat under those circumstances could amount to negligence.

Any process that purports to be based on a rights-based system must not be arbitrary. There should be a clear framework within which decisions are made. This should particularly be so when the essence of the decision is to strip away from the individual the right of autonomous decision making.

OBJECTIVES

What should be the guiding objectives and principles upon which any such decisions are to be based? Further, how is capacity to be assessed and what are the tests that the law will endeavour to apply?

When the citizen turns to the law to find an answer, it soon becomes apparent that in the UK it is a mess. There is not one separate body of law established to deal with elderly incapacitated people. However, what can be determined from the case law and statutes that do exist, is that the law has constructed a number of overriding objectives.

First, there are the means by which the duties that are owed to those who lack capacity are enforced. This can be illustrated in financial and property matters, where the Court of Protection can be invited to intervene^c. A Receiver can be appointed to take control of a person’s financial affairs and the Receiver is answerable to the Court. The great weakness here is that professionals and others who have concerns about the possible exploitation of an incapacitated individual will often know little of this remote and distant Court. It only has one base, and that is in London. What is more, those who are aware that exploitation is

taking place may be the very people who are perpetrating it. Who then will activate the process?

Whilst we have capacity, it is possible to make plans that would result in avoiding the costly involvement of the Court of Protection. It is possible to complete a simple document called an Enduring Power of Attorney. This must be completed whilst a person still has capacity and allows one to appoint the individual(s) who will control one’s financial affairs when and if one loses capacity. The simplicity of the process can allow those determined to abuse to prevail on the vulnerable person, who may even lack the capacity to execute the document. The principle is that, as we have capacity, we have the right to make unwise decisions. When capacity is lost, the Enduring Power of Attorney becomes effective by registering it with the Court of Protection. This is essentially an administrative task.

Second, the law can be invited to consider what is in the “best interest” of the incapacitated person. This can involve the right to be supported and to be legally represented. However, if the criteria for appointing a receiver under the Court of Protection are met, then the incapacitated person cannot instruct a solicitor or bring legal proceedings in his/her own right. This role must be undertaken by a “litigation friend”¹. This objective is further endorsed by the Human Rights Act 1998 which sets out to promote the fundamental rights, freedoms and liberties of UK citizens. Article 6 protects the right to a fair trial. The difficulty arises when seeking a mechanism to achieve this. Yet again, the process of protection is entirely dependent on someone making the link between the incapacitated person and the judicial process.

Third, and perhaps most significantly, is the role of the law to protect individuals from ill-treatment, abuse, neglect and exploitation. The forms that these take are endless and nauseating, as we see in the range of daily media reports illustrating that people know no bounds in gratifying their excesses against vulnerable people, in areas such as money, possessions and physical and sexual abuse. The obvious difficulty in using the criminal justice system is that the abused person may not have sufficient capacity to give evidence. Unless there is separate, corroborative evidence the perpetrator may go unpunished.

PRINCIPLES

The above objectives are not, and in fact cannot be, freestanding. They are guided by a set of prevailing principles that are transferable to the development of the law itself.

In exploring the principles, it should be noted that much research was done by the Law Commission² and the Lord Chancellor’s Department³ in the area of Capacity.

In UK law, all adults are presumed to be competent and as such will have the same rights and opportunities as others. The burden of proof therefore lies on those who seek to assert the citizen’s lack of capacity. Where there is the need to interfere in the lives of those who lack capacity, any such intervention should be the minimum required in the circumstances. Where an individual has been found not to have capacity, nevertheless, he/she should be encouraged to take the kind of decisions that he/she might have taken before capacity was lost. This may sound like an effective set of principles. However, they need to be set in their practical context.

Sadly, it may be in the interests of everyone concerned that the individual should be treated as if he/she were no longer capable of making his/her own decisions. After all, an elderly person may want to make a decision that we might consider unwise, foolish or

^cCourt of Protection is an Office of the Supreme Court and the provisions cited are to be found in s.93 of the Mental Health Act 1983.

even dangerous. That person could be your mum or dad, who may want to give money away or enter into a close relationship with someone many years younger; your inheritance might be threatened. Asserting his/her incapacity could protect your finances. Or a situation might arise where the elderly person does not like the rules and regulations in the nursing home. He/she may find him/herself being treated as a child. "No, you cannot go out; here is your medicine you must take it; it is time for bed now; what we are doing is in your best interests." Who has proved his/her incapacity? Who has even thought about it? Rights can be very uncomfortable.

The courageous professional who correctly asserts that a person has capacity and that he/she may take responsibility for his/her own decisions may be left vulnerable to criticism and even attempts at litigation. Defending the decision when some damage or loss has ensued against an irate family member or community can be an uncomfortable business. This takes knowledgeable and confident professionals.

Therefore, the confidence of professionals and others who are called upon to exercise these judgements is crucial, as is the extent and timing of the intervention. The law has a further role in determining whether and to what extent the intervention by the state has been *proportionate* to the presenting facts and evidence. A proportionate response is an important concept in the Human Rights Act.

Before concluding this section on the role of the law within the context of incapacity, there is an often-overlooked area, namely the role of the lawyer. The role of the lawyer can become confused if the lawyer has not posed to him/herself the vital questions: who is my client?; who am I actually acting for?; to whom do I owe my professional responsibility?; is it the vulnerable elderly person or the family member? The lawyer has to be alert to the situation in which he/she is requested to advise the family who are trying to use the law to exploit the incapacitated individual, e.g. by assisting the family in obtaining an Enduring Power of Attorney (see above) from someone who, had the proper enquiries been made, would have been found to lack capacity (and therefore unable to execute the deed).

WHO LACKS CAPACITY?

Most decisions about capacity are made without any thought at all about objective criteria. Doctors, nurses, social workers, families, or nursing home owners will simply assert it as a fact in order to justify overriding the wishes of the individual.

As indicated above, the Court is the ultimate arbiter of the issue of capacity. However, in those rare cases where it is asked, it is misleading to believe that the judge will arrive at some clear and objective decision when the very criteria for analysis are based partly on the shrouded mystic of clinical analysis and partly on societal attitudes. Essentially there are no clear criteria and it is not good enough simply to assert that each case will be decided on its own facts. The Human Rights Act should cause a shift in emphasis, where the starting point will be the rights of the citizen. Article 6 guarantees the right to a fair trial. The problem may be, who is going to enforce this? The judiciary will have to be alert.

THE BURDEN OF PROOF

When making this decision, the law asserts that a person is presumed at common law to have legal capacity unless it is shown that they do not. However, it is the medical practitioner who may seek to assist or convince the court of the lack of capacity. If such a question comes before the Court, the law at present places the burden of proving such incapacity on the person who asserts it (on

a balance of probabilities, e.g. 51/49). In itself this is worrying, as a diagnosis of incapacity may lead to the loss freedoms, rights and liberties which, our society says in other contexts, namely criminal justice, require that a much higher standard of proof should be reached. Bearing in mind the consequences of a finding of incapacity, surely the standard of proof to rob a person of the control of his/her own affairs should be a much higher one (i.e. beyond all reasonable doubt).

THE HERE AND NOW TEST

An important principle needs to be understood. In assessing a person's capacity, the law will seek to do so at the relevant time in respect of the particular activity/transaction that the person is about to enter into. Anyone assessing an individual's capacity must apply this test. It is possible that a loss of capacity may be transient or episodic. In the early stages of dementia it may be the subject of partial remission.

What is more, a person may both have and lack capacity at the same moment in time. There are differing tests to be applied, e.g. when making a will, entering into a contract, consenting to marriage or sexual activity, making an Enduring Power of Attorney or consenting to medical treatment. When looking at the statutory test of incapacity to be applied when applying to the Court of Protection to appoint a Receiver to manage the property and (financial) affairs of an incapacitated person, it is required that the person must be incapable "by reason of mental disorder". This is defined at Section 1 of the Mental Health Act 1983.

A person may lack the capacity to make a will but possess sufficient capacity to go shopping or vote. Unless it really is the case, we should refrain from defining a person generally as "incapacitated".

There are various ways of approaching the assessment of capacity. The three key methods were usefully outlined in the Law Commission Consultation Paper⁴. The three approaches are:

- *Outcome*. Capacity is determined by the content of the individual's decision.
- *Status test*. May apply in respect of age/diagnosis, etc. without further consideration of the individual's actual competence.
- *Test of understanding*. This concerns an assessment of whether the particular individual is able to make a particular decision at a particular time.

It is the last of these that applies in our common law and which in fact was proposed by the Law Commission in its *New Jurisdiction* publication of 1991 to form the basis of a statutory test.

As can be seen above, a range of different specific tests exists for particular instances, but normally general principles apply which substantively are based upon the individual's understanding, rather than the exercise of his/her judgement. If we were to be judged by the appropriateness of the decisions we make, incapacity would be a far more common concept!

Having noted the very variable application of the framework, it is disturbing to note that little if any research has been undertaken to determine the skills and abilities necessary to establish capacity⁵.

EVIDENCE

Regardless of the above criteria, from an evidential position mental incapacity is a question of fact and there can be no doubt that the correct legal test must be applied. That, as alluded to above, will vary according to the circumstances. The role of the judge within the court proceedings is seeking to determine capacity as a lay person, influenced by personal observations

and on the strength of evidence submitted from doctors and others who know of the individual^d. In the words of His Honor Mr Justice Nicholas Wall:

Expert witnesses need to remember that most judges do not have any more medical expertise than the average intelligent lay person... it is for this reason that they rely upon the integrity of expert witnesses⁶.

THE “NEXT OF KIN”

Our nearest blood relative. Our kind and loving family. Those who are only capable of acting in our best interest. Surely, in the legal lacunae of decision making they are the legal rock on which we can build. If we are incapable of making decisions for ourselves, then surely professionals must turn to them to sanction actions. They know the best way to spend our money, they know when we should be going into a home, they know what is best. Really? Are you sure?

IN WHOSE BEST INTERESTS?

The UK law confers no legal rights on the “next of kin” *per se*. The matrix of decision making for a person who lacks capacity is our common law doctrine of necessity. This involves four basic elements:

- The person must lack capacity in relation to that particular decision at that time.
- It must be sufficiently necessary to make that decision.
- The person making it must be reasonable.
- It must be in the best interests of the incapacitated person (not the next of kin and not the professional).

All too often the professionals may feel powerless in the face of families who are demanding admission to a nursing home, or a discharge from hospital. They may be trying to insist on a particular form of treatment or withholding their consent to what the medical team is proposing. The professionals know that no “reasonable person” could possibly believe that certain decisions were in the incapacitated person’s best interests. But, the family want it, so how can I possibly prevent it? The lawyer would point out that if the professionals capitulate and damage ensues, that it could be called professional negligence.

TREATMENT

To have capacity gives us the right to refuse any treatment that we do not wish to receive, even if death would be the result. Only in the case of people detained under the Mental Health Act can a competent adult be treated against his/her wishes, and then only for his/her mental disorder.

Clearly drawn and unambiguous “advance refusals” (otherwise known as living wills or advance directives) allow persons with capacity to bind the hands of the doctors. They can indicate that in the event of them becoming incapacitated, there would be certain treatments that, had they retained capacity, they would not have consented to. These have emerged through our case law, notably the case of *Airedale NHS Trust v. Bland* [1993] 1 All E.R. 821 and are a common law concept. An advance refusal could only be overridden if either it was not sufficiently clear or the

person was then “sectioned” under the Mental Health Act because he/she required treatment in a hospital for mental disorder. Our statute law always takes precedence over common law principles.

WHEN TO “SECTION” UNDER THE MENTAL HEALTH ACT

From my own experience of training mental health professionals for many years, few professionals working with the elderly seem to realize that the Mental Health Act provides a framework of protective powers for those who lack capacity and have a mental disorder. This term includes the mentally ill. Dementia is of course a form of mental illness.

The Act allows a person to be “sectioned” when his/her mental disorder is of a nature or degree that requires either assessment (section 2) or treatment (section 3) in hospital and his/her health, safety or the protection of others require it. Once sectioned it allows the professionals to impose their will on the patient but certain boundaries are proscribed. It gives the individual certain rights of appeal, and these can be triggered simply on the basis that the right has not been exercised.

Since the House of Lords judgement in the case of *R v. Bournemouth Community and Mental Health NHS Trust ex parte L* 1998 3 W.L.R. 107, the incompetent patient need only be sectioned if it is necessary to treat him/her in hospital for a mental disorder and he/she demonstrates actual objection by word or deed. Because statute law (that passed by Parliament) takes precedence over our common law, the person should be sectioned, and if he/she is not, then any treatment given under the common law doctrine of necessity would amount to unlawful imprisonment and assault. You cannot use the defences available under common law if you are obliged to apply the statute law. At present, it would seem to me that few medical and nursing professionals realize this. The guidance issued by the Department of Health⁷ following the Bournemouth judgement tells us that if there is doubt as to whether the action amounts to an objection, then it is to be treated as if it were an objection. The incapacitated person who is trying to leave the ward or is objecting to medication (e.g. spitting it out), where there is doubt, should be considered for a Mental Health Act section. This would then accord him/her rights under that Act. It would then follow that further treatment for mental disorder under common law would be unlawful.

What about the person who is “required” to go into a nursing or residential home and does not wish to go? The Code of Practice to the Mental Health Act, Chapter 13 paragraph 13.10.b., indicates that, under these circumstances, Guardianship under the statutory powers in the Mental Health Act should be seriously considered.

Where an adult is assessed as requiring residential care, but owing to mental incapacity is unable to make a decision as to whether he/she wishes to be placed in residential care, those who are responsible for his/her care should consider the applicability and appropriateness of guardianship for providing the framework within which decisions about his/her current and future care can be planned.

Few professionals seem to realize this.

A Guardianship Order allows a guardian to require a person to “reside in a particular place”, attend various places for occupation, education or treatment or to allow access to the person on Guardianship by specified third parties or the guardian.

^dThis proposition is strongly supported in the High Court Judgement of Wall J in *Re: G (Minors) (Expert Witnesses)* [1994] 2 FLR 291, 298.

Although the use of Guardianship is increasing with the elderly (Mental Health Act Commission 7th Biennial Report⁸), it is still used infrequently. There were 804 Guardianship cases current at 31 March 1998 compared with 335 in 1992. This, however, includes all age groups and mental disorders. There are no statistics available specifically for the over-65s. Usually the common law doctrine of necessity is invoked and this allows reasonable force to be used where necessary, either to prevent a person leaving or to restrain him/her for medication or other reasons. This is likely to be subject to challenge under the Human Rights Act 1998. There is a strong possibility that the Bourne-wood case will be overturned by a decision of the European Court of Human Rights. This is because, under Article 5, there is a Right to Liberty and Security of the Person. This states under Article 5(4) that:

Everyone who is deprived of his liberty . . . shall be entitled to take proceedings by which the lawfulness of his detention shall be decided speedily by a court and his release ordered if the detention is not lawful.

For those who lack capacity and are “detained” under common law there is not a mechanism that allows this challenge in English law.

THE NEW ERA OF HUMAN RIGHTS

With the passage of the Human Rights Act 1998 we are entering a new era for the concept of human rights in the UK. With the partial incorporation of the European Convention of Human Rights and Fundamental Freedoms there is the opportunity for the citizen to take proceedings against public bodies for alleged breaches of human rights. Public Bodies would include state and private hospitals, social service authorities, doctors and social workers who can carry out specific functions under the Mental Health Act where any of these bodies are carrying out a public function. As appears usual in the legal process, the mechanism by which the incapacitated can seek to assert their rights is far from clear. Yet again it will require the identification of the individual who will take the proceedings on their behalf.

The most relevant Articles to the European Convention of Human Rights and Fundamental Freedoms are:

- Article 2 enshrines the right to life. This includes the right to be protected by those who are caring for the vulnerable from risks that may lead to death.
- Article 3 protects the citizen from torture, inhuman or degrading treatment or punishment.
- Article 5 states that there is a Right to Liberty and Security of the Person.
- Article 6 protects the right to a fair trial when, for example, the Court of Protection is considering removing a person’s right to control his/her own financial affairs.
- Article 8 provides for respect for a person’s home and family life. This would include decisions about confidentiality of information, where a person lives or with whom he/she has contact.
- Article 12 provides for the right to marry (despite one’s age!).

IN CONCLUSION

With an inadequate legal framework, extensive abuse taking place in many guises, with those working with the elderly often unaware of what the law permits, with the government apparently unwilling to legislate, the future looks bleak. The Human Rights Act could and should be about changing our culture. It should affect the way that we conceptualize the exercise of decisions that affect the rights of the citizen. My fear is that in all other areas of society this may well be the case. However, for the elderly confused incapacitated person, who will take on responsibility for this awesome responsibility?

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Older People, Clinicians and Mental Health Regulation

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The Mental Health Act 1983 (England and Wales) made no mention of age, applying to children and those of advanced years just the same as to younger adults. Similarly, the Mental Health Act Commission created by the 1983 Act to “keep under review the operation of the Act” as it related to detained patients also had no responsibilities specific to older people. The old Commission had an awkwardly circumscribed role with few real powers. It visited hospitals and registered nursing homes to meet patients and ensure that the conditions in which they were detained were of an acceptable standard. The Commission also investigated complaints relating to detained patients and monitored the consent to treatment safeguards in Part IV of the Act, appointing independent doctors to give second opinions.

In December 2000, after 10 years of discussion and consultation about the failings of the 1983 Act, the government published proposals for a new Mental Health Act¹, which will have a very significant impact on the practice of geriatric psychiatrists. It contains new safeguards for the care and treatment of people with long-term mental disorders in institutional care, providing obligatory second opinions for those who are unable, through mental incapacity, to consent to treatment. In effect, this will cover all those with dementia in long-term care and many other older psychiatric patients. The new Act also replaces the old Commission with a new Commission for Mental Health, which will have wide powers in relation to overseeing the new legislation but will not be a visitorial or inspectorial body. For the first time, the new Commission will have a responsibility to ensure that professionals are trained properly in the legislation. This will surely require far closer working relationships with professional training and accreditation agencies. The old visitorial function, however, will be handed over to the new National Care Standards Commission and the Commission for Health Improvement. Thus, clinicians can expect even more inspection and regulation, rather than less, under the new regulations.

The central regulatory system is usually dated back to 1833, when the Factory Inspectorate was established. *The Times*² pronounced that this new system contained “the seeds of mighty changes”, although the Editor was “no enthusiast” for central regulation but acknowledged that an inspectorate offered advantages “if inspectors or visitors of strong capacity, of enlightened humanity and moral courage” were appointed. The Lunacy Commission of 1845, chaired by the indefatigable 7th Earl of Shaftesbury for 40 years until his death in office, achieved considerable influence with government and changes in local asylums and workhouses because the Commission remained

small, elite and adopted a coherent, unifying set of policies in its early years³.

Modern mental health commissions are similar to the Lunacy Commission in being only as effective as their members are. Ministers have not always been convinced of this simple truth and have sometimes seen Commission appointments as a convenient reward for other fields of endeavour or as an opportunity to promote other laudable government objectives. Since its inception in 1984, the old Commission, a multiprofessional body, struggled to attract members of distinction from the professions of psychiatry and law and yet, to achieve credibility and respect from psychiatric services, the quality, training and behaviour of members was crucial. Over the years there was a steady improvement in the administrative efficiency of the organization and a significant step up in quality of the recruits. The new Commission will need to learn some of the lessons learnt if it is to achieve early credibility. Being a good commissioner requires enormous tact, humility and an ever-present awareness that a nurturing, developmental, encouraging approach achieves far more and is less alienating to professional staff than a heavy-handed “policing” approach.

The notion underpinning regulatory bodies dies hard. The idea is that the Secretary of State employs a team of quasi-independent “eyes and ears” to act as the conduit for information to central government and to channel edicts from the centre to the field. The proliferation of statutory commissions and non-statutory regulatory bodies (so-called QUANGOS) in the late twentieth and early twenty-first centuries would suggest that faith in these institutions as movers and shakers of social improvement remains undimmed in governments today. The zeal with which agencies are established falls away as soon as it is realized that inspectors and monitors cannot substitute for good local managers. In mental health services, good hospital unit management and improvements of standards of training and clinical work through professional bodies, such as the Royal College of Psychiatrists and the National Boards for nursing education, are more likely to effect permanent improvements in standards of care. “Watch-dogs” and Commissions inevitably disappoint ministers and the usual cycle of events is that a commission’s powers are progressively reduced and in due course frequently disbanded on the grounds of economy. As the numbers of factories grew beyond what it was reasonable to inspect, the Factory Inspectorate’s sweeping powers to make statutory regulations and act as local magistrates were abolished in 1844. The Board of Control similarly found its powers diminished from those of its predecessor, the old “dead duck” Lunacy Commission, and was

finally abolished in 1959. More recently, the Health Advisory Service, established by Richard Crossman's health ministry in the 1960s, was perceived to be unable to stop hospital scandals and was gradually denuded of its powers and eventually extruded from agency status to make a living from consultancy as best it could. Now the Mental Health Act Commission predictably gives way to a new-style Commission, with more circumscribed, focused powers, and it remains to be seen whether the new one will fare any better in the eyes of services and the government than the old one.

It has been estimated that there are approximately 44 000 informal admissions to institutions annually in England and Wales of mentally incapacitated patients who are compliant with treatment but lack the capacity to consent to treatment⁴. This far exceeds the 13 000 detained under formal powers. The new Act will provide them with extra safeguards, the right to a second opinion for long-term treatment and the right to appeal to a Tribunal. Some psychiatrists will regard an extension of legal powers to informal incapacitated patients as an unwelcome extra burden of work on them and their clinical teams. On the other hand, the new provisions will ensure that older people will receive greater attention from the regulatory bodies and their legal rights to decent care and treatment will be enhanced.

When they work well, central regulatory Mental Health Commissions can be strong allies to clinicians seeking to improve their patients' lives. Psychiatrists need to understand the role and remit of the Commission and be willing to serve during part of their career. The effectiveness of the new Commission will be greatly enhanced if the profession adopts a strategy of supporting and involving itself in its work and also ensures that the needs of older people are kept firmly in the forefront of the regulators' considerations.

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Training Requirements for Old Age Psychiatrists in the UK

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In the UK over the last 20 years, increasing numbers of psychiatrists have specialized in working with older adults. Various terms have been used to describe this area of work: “psychogeriatrics”, “geriatric psychiatry” and “old age psychiatry” are probably the most common. In 1978 the Royal College of Psychiatrists formed a Section (now Faculty) of Old Age Psychiatry and just over 10 years later, in 1989, old age psychiatry became recognized as a specialty by the Department of Health. The Royal Colleges of Physicians and Psychiatrists produced a Joint Report in 1989, which devoted a chapter to education and training in the psychiatry of old age¹. At that time there was no accepted training programme for psychiatrists aiming for a career in the developing specialty. Things have changed considerably over recent years.

BACKGROUND: GROWTH OF THE SPECIALTY

In the early 1980s, two surveys of psychogeriatric services^{2,3} showed considerable growth in the developing specialty, such that over 200 consultants were identified by late 1983. At that time the Joint Committee on Higher Psychiatric Training of the Royal College of Psychiatrists required higher trainees to spend 12–18 months in posts where the bulk of the work was in the subject; 28 senior registrar placements were identified. The authors predicted ongoing growth in old age psychiatry and a need to expand available training.

By 1990 there were 360 consultants working mainly in old age psychiatry and in 1993 the total had increased to 405⁴. Figures collected by the Faculty of Old Age Psychiatry show that the total has continued to rise, although there is still a high vacancy rate of approximately 14%.

UNDERGRADUATE EDUCATION

The first Joint Report¹ recommended that each medical school should have a senior academic in old age psychiatry and that all medical undergraduates should receive training in the subject. Faire and Katona⁵ surveyed undergraduate teaching in the UK and reported a considerable expansion of academic posts in the specialty, but noted that more than half of all departments lacked a senior old age psychiatry academic. Almost all medical schools offered formal lectures in the subject, but there was great variation in the amount of clinical experience on offer and the authors felt that there was a strong case for all medical students having clinical

experience in old age psychiatry, as recommended by the Joint Report. Gregson and Denning⁶ surveyed teaching hospital psychiatrists and found that many teachers set no formal learning objectives in old age psychiatry. Most respondents wanted their teaching to impart enthusiasm for the subject, a sense of hope in working with mentally ill older adults, and an awareness of issues specific to ageing and ageism.

Although there are now chairs or readerships at a number of medical schools in the UK, gaps remain⁷ and the second Joint Report, published in 1998, recommends that the characteristics of mental disorders among older people and the principles of good quality care should be included in the core curricula of all schools of medicine and nursing.

POSTGRADUATE TRAINING

The total minimum duration of specialist training in old age psychiatry is 6 years, of which 3 years will be in general professional or basic specialist training and 3 years in higher training (as a specialist registrar).

Basic Specialist Training

Part II of the Membership examination of the Royal College of Psychiatrists (MRCPsych) can normally only be taken after 30 months of training in psychiatry and is a requirement for entry into higher training, so basic training normally lasts for about 3 years. The first 12 months may include 6 months in old age psychiatry, provided that the experience offered is broad and includes the assessment and treatment of people with functional mental illness. Experience in old age psychiatry during basic training is regarded as important because of the increasing elderly population and the high rate of mental illness in older people, but the College *Basic Specialist Training Handbook*⁸ states that trainees should be exposed to acute and functional mental disorders in late life and not solely to organic brain diseases. Old age psychiatry placements can often offer good community experience for trainees and the opportunity to attract young psychiatrists into the specialty.

Basic training concentrates on providing a range of experience in the specialties and subspecialties of psychiatry, aiming to develop history taking, formulation and case presentation skills, therapeutic skills and clinical judgement, relationships with

colleagues, patients and relatives/carers, basic psychiatric knowledge and appropriate knowledge of general medicine.

By the time the trainee is ready to move into higher training, he/she will have completed a minimum of 3 years in approved training placements and will hold the MRCPsych⁹. He/she will also have some idea of his/her eventual career intentions. General psychiatry and old age psychiatry Specialist Registrar posts may be advertised separately, but trainees may opt to undertake training jointly in both specialties (see below).

Higher Specialist Training

Higher training aims to provide an educational programme to prepare a trainee for independent practice in old age psychiatry. The number of higher trainees is determined by the number of national training numbers (NTNs) and this is fixed centrally by the NHS Management Executive or equivalent body. The Specialist Training Committee of the Royal College of Psychiatrists (STC) sets the standard for training schemes and, under the aegis of the Specialist Training Authority (STA), sets the standard for award of certificates of completion of specialist training (CCSTs), which indicate that specialist training has been successfully completed. Since 1997 a CCST has been mandatory before taking up an NHS consultant post. Old age psychiatry falls within the remit of the Royal College of Psychiatrists' General and Old Age Psychiatry Specialist Advisory Committee (GOAPSAC)¹⁰.

General and old age psychiatry higher training schemes may offer two options for aspiring old age psychiatrists. Single accreditation involves training for 3 years to gain a CCST in old age psychiatry. Currently these trainees may spend 1 year in general psychiatry or one of its subspecialties if they so wish. Single accreditation therefore necessitates a total of at least 6 years in psychiatric training. Many trainees (probably about 60%) opt to complete dual training, which aims at dual certification in general and old age psychiatry. These trainees complete a 4 year higher training programme, with 2 years in each specialty. They must have been appointed to their specialist registrar posts by an appropriately constituted appointments committee and will hold a NTN in old age psychiatry. Dual certification will require a total of at least 7 years in psychiatric training.

During their training, specialist registrars are expected to develop their professional attributes, core knowledge and skills, and are set goals in research and audit, teaching and supervision, and management. Currently, six "core" sessions are devoted to experience in old age psychiatry (or other specialty). "Core" experience involves working with a multidisciplinary team to provide a service to a defined population. Two further sessions are available for research, audit and personal study, and another two can be used to develop special interests. Old age psychiatry trainees are expected to gain experience of geriatric medicine at some stage of their training, and this is usually achieved either on a short-term attachment or using special interest sessions.

CONTINUING PROFESSIONAL DEVELOPMENT

Loane and Barker¹¹ surveyed newly appointed old age psychiatrists' views of their higher training. Overall clinical experience was felt to be satisfactory, but management experience was lacking in a number of areas and experience in dealing with complaints, dealing with difficult professional relationships, recruitment and disciplinary proceedings were all identified as areas where training was insufficient. Higher trainees are expected

to get training in management but it can be difficult to pitch it at the right level.

The emphasis today is on lifelong learning¹², which is regarded as essential for all healthcare professionals. Old age psychiatrists are no different and are likely to see continuing developments in their field throughout their working lifetimes. Learning does not stop at the transition from higher trainee to consultant, and some might say that this is the point at which learning really starts. Consultants increasingly plan their CPD programmes¹³, although these need to be flexible and to evolve with the specialty, the individual and the job. Increasingly too, consultants change their interests, disciplines and posts as their careers progress. This may be a way to re-energize and deal with the stresses of their multiple roles¹⁴. Continuing professional development should be a positive supportive opportunity for consultants to continue learning throughout their working lives.

LIFELONG LEARNING

Learning about old age psychiatry starts in medical school and continues throughout the working life of an old age psychiatrist. The context within which the specialty operates is constantly changing. There are various threats and opportunities on the horizon, including changes to the Mental Health Act¹⁵, new ways of dealing with people unable to consent¹⁶ and the National Service Framework for older adults. Old Age Psychiatry and its practitioners cannot stand still. The enthusiasm which teaching hospital psychiatrists aim to impart to their medical undergraduates can be maintained during specialist training and boosted throughout a consultant's career by continuing professional development and the challenge of working within a constantly changing health and social service context.

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Old Age Psychiatrists and Stress

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Stress, and its effect on the workforce, is a matter of increasing concern in the Health Service. Stress levels in health professionals generally are high¹. Doctors as a group have increased rates of cirrhosis, road traffic accidents and suicide, compared with the general population². They are prone to symptoms of anxiety and depression, and are more likely to misuse alcohol or other substances³.

WHY WORRY ABOUT SERVICES FOR OLDER PEOPLE?

Working with older people, especially those with mental health problems, might be particularly stressful, for various reasons. This client group is more likely to exhibit challenging behaviours. They are subject to the increasing disadvantages, disabilities and progressive loss of independence associated with increasing age, and are approaching death. Consultants (and other staff) in geriatric medicine and geriatric psychiatry are working in so-called "Cinderella specialties", which struggle to compete for resources with the more "sexy" acute specialties. The stigma of being old, mentally ill and cognitively impaired is contagious and affects attitudes towards the staff who work in these specialties⁴. In addition, staff will have to confront their own beliefs and fears about ageing, dementia and death for themselves and members of their own families⁵.

It is not surprising, in this context, that psychiatrists are retiring earlier⁶ and recruitment to the specialty is inadequate to maintain consultant numbers⁷. Stress is an important issue for the workforce.

WHAT DO WE KNOW?

Studies of the work patterns of old age psychiatrists have found that they have long working days with little opportunity for recreation, family life, personal study and research^{8,9}. More than 40% of old age psychiatrists do extra work at home on every day of the working week except Friday, and more than 30% do so on Saturdays and Sundays⁹. Most of the stresses identified by old age psychiatrists relate to work overload or organizational structure and climate¹⁰. Many of these factors are equally applicable to staff working in geriatric medicine.

WHAT CAN BE DONE?

Appointment as a consultant brings long working hours and a number of different, often conflicting, roles (including responsible clinician, manager, budget holder, counsellor, researcher, teacher, team member, perhaps team leader) in various settings (wards, day hospitals, community and others). The result is role ambiguity, conflict and overload. Doctors could be better trained for the demands of consultanthood. The means by which consultants are supported, supervised and valued could be radically revised¹¹. Individuals need to be able to change and develop their interests and work patterns over time, in order to allow re-energization. Time allocated to clinical work, teaching, research, family and other interests will vary at different stages of a person's working life. Organizations need to accept and support the evolving careers of their staff members.

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Developing and Maintaining Links between Service Disciplines: the Program for Organizing Interdisciplinary Self-education (POISE)

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The need for effective and systematic education of geriatric specialists in the field of psychiatry has been widely recognized¹. Current shortages of general and child psychiatrists are gradually being surpassed by the shortages of geriatric psychiatrists². The field of geriatric psychiatry education began to expand only recently, and this expansion was a direct result of funding of postgraduate specialty training programs in geriatric mental health by the National Institute of Mental Health³.

Another major development in the field of psychiatry that has led to increased interest in, and expansion of, geriatric psychiatry training programs has been the evolution of subspecialization in geriatric psychiatry through Added Qualifications (Board Examinations) and the accreditation of geriatric psychiatry residency/fellowship programs. However, as the specialty field of geriatric psychiatry has evolved, it has become more apparent than ever that general practitioners, rather than geriatric psychiatrists, will provide the bulk of care to mentally ill older people. Although guidelines for developing curricula in geriatric psychiatry have been developed^{1,3}, these guidelines focus primarily on the clinical skills necessary for the psychiatrist or general practitioner to treat geriatric patients, and not on the leadership skills necessary for the psychiatrist or general practitioner to facilitate and lead the interdisciplinary healthcare team. The curriculum guidelines also neglect to emphasize the role of the geriatric psychiatrist as the key link between service disciplines on the mental healthcare team.

For some time, the interdisciplinary healthcare team approach has been well established in specialty areas, such as rehabilitation, surgery and dentistry; however, the fields of psychiatry and general medicine, specifically geriatric psychiatry, have been slow to realize the important role of the geriatric psychiatrist as part of the mental healthcare team approach to care of the geriatric patient. The field has been even slower to recognize the central role the geriatric psychiatrist can and should play in facilitating cohesive team function and linking treatment goals of different service disciplines. The purpose of this chapter is to describe a model interdisciplinary team leadership training program, which exists within the curriculum of a geriatric psychiatry fellowship program sponsored by the New York State Office of Mental Health. This model program is unique in that it focuses on training geriatric psychiatrists in the skills required to develop, lead and maintain interdisciplinary treatment teams in inpatient and outpatient settings.

Schmitt *et al.*⁴ indicate that the term “interdisciplinary teams” in healthcare settings has a variety of meanings, depending on

usage. Thus, it is important to establish criteria that define the term. For the purpose of this chapter, we have adopted the criteria set out by Schmitt and her colleagues⁵. These criteria require at a minimum that the healthcare team: (a) includes a variety of disciplines in the care of the same patient; (b) encompasses a diversity of dissimilar knowledge and skills required to treat the patient; (c) plans care by establishing an integrated set of goals shared by the providers of that care; and (d) shares information and coordinates their services through a systematic communication process.

PROGRAM BACKGROUND

The interdisciplinary team training program described in this chapter is the outcome of 15 years of work devoted to the development of a durable, cost-effective method of linking and coordinating mental health services within the institutional setting. The program evolved, in part, from an educational philosophy that focuses on a participative model of self-education. The assumption is that the healthcare staff of an institution already have the technical skills required to function in their particular position, but need to enhance their understanding of how their roles in their particular discipline relate to the roles of other team members from different disciplines, and how the team as a whole relates to other staff and teams. A system by which the staff can work collaboratively to deliver effective treatment to patients is regarded as essential.

This concept was first applied in assessment training programs and subsequent related studies involving mental healthcare teams at state psychiatric centers in New York⁶⁻⁸. The results of these evaluation studies and the favorable response of staff to the interdisciplinary mental healthcare team leadership training⁹⁻¹¹ led to a request by the Deputy Commissioner of the New York State Office of Mental Health to adapt the program for implementation in other psychiatric centers throughout New York State. In subsequent discussions, it was determined that the training would be most successful and relevant if it was conducted with the key member of the mental healthcare team who most often is responsible for team leadership, namely the psychiatrist. In this way, a culture of learning would be established and the psychiatrist trainees would then go on to work on interdisciplinary treatment teams elsewhere, and bring with them to their new work setting this culture of learning. Thus, the psychiatrist trainees would

disseminate to other staff members from other disciplines the knowledge they obtained during the team leadership training. This adaptation of the original model program was applied to the development of the Columbia University Geriatric Psychiatry Residency and Fellowship Programs, which are sponsored by the Stroud Center and the Department of Psychiatry of the Columbia University Faculty of Medicine, Binghamton Psychiatric Center of the New York State Office of Mental Health. This program is accredited by the Accreditation Council for Graduate Medical Education. The Geriatric Psychiatry Residency and Fellowship Programs bring together the clinical resources of the Binghamton Psychiatric Center and the educational and clinical research resources of the Stroud Center/Center for Geriatrics and Gerontology of Columbia University.

Since its inception, the Geriatric Psychiatry Residency and Fellowship Programs has included, as a core ingredient of the long-term care component of the curriculum¹², the training of psychiatrist-fellows in the methods of interdisciplinary team leadership. Additionally, every fellow is required, during his/her fellowship, to develop and/or lead an interdisciplinary treatment team under direct and regular supervision of fellowship faculty.

PROGRAM DEVELOPMENT

Rationale

All clinicians need a good system for identifying patients' symptoms, making informed treatment decisions regarding the patients, and managing stress that is related to providing care. All members of the treatment team need practical tools to guide them in eliciting, classifying, recording, and interpreting information on patients' health status and functioning—in short, a system that will help them to evaluate each patient's status, identify the appropriate treatment, predict possible outcomes, and plan the patient's care. This system must also include methods to assist the interdisciplinary team in functioning as a team and managing their own stress.

A program designed to train interdisciplinary mental health care team members at psychiatric centers within the New York State Office of Mental Health was developed, implemented and evaluated^{8,10}. The purpose of the program was to upgrade the functioning of the multidisciplinary/interdisciplinary mental healthcare treatment team and to train staff to identify and manage the stress that developed as a result of caring for older psychiatric patients, many of whom are demented. The Program for Organizing Interdisciplinary Self-education (POISE), the name by which the program is officially known, is a multidisciplinary/interdisciplinary approach to improving the assessment of patients in psychiatric hospitals and the treatment planning decisions based on those assessments. The program also provides training of staff in current methods of stress management. POISE currently focuses on the training of the geriatric psychiatrist as the central member of the interdisciplinary mental healthcare team.

The geriatric psychiatrist is a key figure in the treatment and management of geriatric patients in both inpatient and outpatient psychiatric facilities. The geriatric psychiatrist is also a core member of the interdisciplinary treatment team and is a key link between service disciplines on the team in psychiatric settings. Regardless of the level of functioning of the team, the geriatric psychiatrist is often viewed by team members as the primary care physician who is ultimately responsible for leading the team and thus coordinating the treatment of geriatric patients. However, most geriatric psychiatrists receive little or no training that will enable them to work effectively as a core member of the interdisciplinary team and key link between team members.

The geriatric psychiatrist and other members of the interdisciplinary team require special training to enable them to work effectively together^{13,16}. Training, which can be accomplished either formally or informally on and/or off the unit, can be directed at facilitating and encouraging a team approach to patient care. This training must be durable in as much as it can be replicated throughout an entire system of care to achieve objectives; in that its effects last beyond the period of training; in that it concentrates on developing team cohesiveness and increased productivity; and in as much as it is based on teaching the team how to continue and maintain self-learning processes (including monitoring of own performance to achieve objectives). The training must be directed at a key member of the team (e.g. geriatric psychiatrist), who then goes on to train other members of the team. In this way, team functioning can continue, even in the absence of the facilitator, the geriatric psychiatrist.

The Columbia University Geriatric Psychiatry Residency and Fellowship Programs includes training in interdisciplinary team leadership as a core component of its training of geriatric psychiatrists. This training component, POISE, is described below. Details regarding the specific interdisciplinary treatment team training approaches used in the training of geriatric psychiatrists are given in Toner *et al.*¹⁴ and Miller and Toner¹⁵, who have outlined the steps in developing and implementing a geriatric team.

Program Description

POISE is a durable, cost-effective approach to teaching geriatric psychiatrists (i.e. residents and fellows in training) methods of self-learning which serve as ongoing tools for planning treatment for patients. It is durable and cost-effective, in as much as the geriatric psychiatrist, and ultimately other members of the interdisciplinary team, learn methods of effectively working with one another by collaboratively setting goals and arriving at appropriate treatment decisions for the patient. After the geriatric psychiatry resident/fellow receives core training in interdisciplinary team development and leadership, he/she is assigned to an existing interdisciplinary team, where he/she imparts the core training to the members of the team. The team then continues to apply these methods on an ongoing basis and the geriatric psychiatrist continues to provide guidance and leadership as the group leader/facilitator. This approach to staff training is untraditional because, instead of relying on conventional didactic approaches to learning or the charismatic qualities of the group leader, the core training focuses on teaching the geriatric psychiatrist the methods of self-learning in regard to assessment¹⁶. Geriatric assessment serves as the unifying theme of the training, because most staff conduct assessment in one way or another—including the nursing aides, who generally do not view themselves as assessors—and because most staff feel they need additional training in assessment. Furthermore, although most team members assess patient functioning, no single discipline considers assessment as their exclusive domain. In this way, while team members are organized and ultimately conditioned to focus on the theme of assessment, interdisciplinary conflict revolving around disciplinary territorial issues is avoided, since no one discipline or individual has exclusive rights to the theme. The theme of assessment also serves as a springboard for discussing more general group functioning issues.

In POISE, the concept of self-learning is applied directly to the training of the geriatric psychiatrist resident/fellow in a long-term care setting serving the elderly. Thus, the geriatric psychiatrist resident/fellow and other members of the interdisciplinary team are trained to develop their own skills and strengths in regard to the identification, classification and

treatment of patient problems, rather than superimposing a costly didactic approach which, at best, can expect to yield only short-term effects because of frequent staff turnover and administrative changes.

More concretely, the geriatric psychiatrist is trained in the methods of team leadership, using geriatric assessment as a central theme, and the methods of facilitating interdisciplinary team collaboration among the kinds of people associated with the chain of patient care, the treatment team. The geriatric psychiatrist also receives, as part of his/her core training, instruction in a method of developing with the team the universe of management decisions available to them in that particular setting (i.e. the Treatment Decision Guide). By using case examples, review of patient records, videotapes of case conferences, admission interviews, etc., the geriatric psychiatrist (as facilitator) and the team arrive at an understanding of good and bad management decisions, what problems exist in making treatment decisions, how these problems develop, what treatment options are (or might be) available, what methods team members use in arriving at appropriate treatment decisions, and what possible alternatives exist in arriving at a diagnosis. By establishing with the team the criteria that must be met in order to arrive at a particular treatment decision, the geriatric psychiatrist and other team members establish goals and objectives for any specific treatment decision available to them and discuss how assessment can be used to better link patient problems to the appropriate treatment decision. Through this process, the Treatment Decision Guide^{14,15} is developed. This Treatment Decision Guide identifies the dynamic pathway for use of information regarding the patient in planning, implementing and monitoring treatment. The Treatment Decision Guide provides team members with a guide for using assessment and is also a useful, cost-effective and durable recipe for planning treatment.

Clinical Applications

POISE sensitizes the geriatric psychiatrist to the psychosocial problems that have a high frequency in the elderly long-term care patient population. It provides the geriatric psychiatrist with a program for upgrading the interaction process between interdisciplinary team members through group process exercises and lectures, which emphasize the role of the geriatric psychiatrist and other team members in assessing the well-being of patients and linking that assessment to treatment. The program then goes on to train the geriatric psychiatrist to lead teams and train other team members in team leadership.

POISE also improves the quality of staff interaction and team functioning by: (a) setting up systematic approaches and procedures for making appropriate treatment decisions; and (b) once this system has been set up, providing a device by which professionals and paraprofessionals relate to one another in operating the ongoing system themselves. This is done primarily through the use of the Treatment Decision Guide and Systematic Problem Solving. In addition, POISE enhances patient care by improving staff knowledge and skills in regard to assessment. This is accomplished through group exercises designed to build the geriatric psychiatrist's and other team members' skills in observing patients and formulating decision plans based on those observations. In this way, the geriatric psychiatrist is trained to be a facilitator, who provides the team with a mechanism by which the group does its own learning on an ongoing basis.

SUMMARY

The Program for Organizing Interdisciplinary Self-education, POISE¹⁶, is one program that has demonstrated that the

principles of interdisciplinary collaboration and team training can be applied successfully in a geriatric psychiatry residency/fellowship program¹³. The learning process involved in POISE covers the skills, knowledge and attitudes of team members regarding the following program components: team development, management, and maintenance; and program needs assessment.

POISE focuses on the training of geriatric psychiatrist fellows in the following areas¹³:

1. The role each member of the team plays in assessing the patient.
2. Linking each member's assessment to a treatment plan.
3. Systematic approaches and procedures for making appropriate treatment decisions. This includes methods of defining and negotiating team members' roles; case study approaches to prioritizing treatment goals; and systematic approaches to problem solving.
4. The Treatment Decision Guide (TDG): a guide designed by the interdisciplinary team members specifically for patient management in their particular institution, the TDG provides a key to arriving at available treatment alternatives in that institution.

After the geriatric psychiatrist is trained in POISE, he/she is trained in the method of teaching other interdisciplinary treatment team members the POISE method and thus provides the team members with the crucial techniques for maintaining the ongoing team themselves. Once the geriatric psychiatrist is assigned to the interdisciplinary team, the emphasis of the training is on shared leadership functions of all team members, with designated leaders, experiential learning to facilitate team functioning and problem solving, and ongoing orientation of new team members (to team objectives and norms).

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Appendix— International Psychogeriatric Association (IPA)

Barry Reisberg (former IPA President) and Fern F. Finkel (Executive Director)

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The International Psychogeriatric Association is a worldwide, multidisciplinary group of healthcare professionals dedicated to the advancement of mental health in the elderly and the field of psychogeriatrics.

The organization was proposed by Imre Fejer and Hans Reichenfeld in 1980, following a very popular course in psychogeriatrics that had been developed by Professor Tom Arie in Nottingham, UK. The organization's founding meeting was held in Cairo, Egypt, in November 1982. Subsequently, IPA congresses have been held every 2 years. Early congresses were held in Umeå, Sweden (1985), Chicago, USA (1987), Tokyo, Japan (1989) and Rome, Italy (1991). The organization has maintained these widely geographically dispersed settings for its congresses until the present time (Table 142.1). In its early years, the growth of the organization was shepherded by Manfred Bergener (Cologne, Germany), President and Sanford Finkel (Chicago, USA), Secretary-Treasurer. A measure of the influence that Sanford Finkel has had on the organization includes his unique role in having served as IPA President (Table 142.2), as well as Secretary-Treasurer and chair of many important IPA committees and task forces.

From the outset, IPA has been successful in contributing to major developments in the field of psychogeriatrics. The organization has done this in large part through its very successful conferences. IPA's Board of Directors has also contributed very significantly. Currently, there are 25 directors and five officers from around the world serving on IPA's Board. From the very beginning, the Board decided that no more than two directors can represent any single country. Consequently, IPA's Board is a disparate and geographically diverse group. Many of the outstanding scientific and medical leaders in our field have served on IPA's board. Examples include Sir Martin Roth (UK), Luigi Amaducci (Italy) and Kazuo Hasegawa (Japan). IPA has also, from the outset, had regularly scheduled regional meetings at intervals of approximately 6 months. These meetings have also been held in diverse locations around the world (Table 142.1). A great advantage of these meetings is that, unlike the IPA congresses, they are generally held in conjunction with a local organization with a neuropsychogeriatric interest. This has enabled IPA to have fertile interchanges with many local professional groups. These include joint meetings with ongoing local or regional groups, such as the Turkish Society of Psychogeriatrics, the Institute of Mental Health of Beijing Medical University, the Royal College of Psychiatrists' Faculty of Old Age and the Brazilian Association of Geriatric Neuropsychiatry.

Table 142.1 IPA congress and meeting dates and locations

November 22–25, 1982	1st Congress; Cairo, Egypt
October 18–20, 1984	Cologne, Germany
August 28–31, 1985	2nd Congress; Umeå, Sweden
September 5–6, 1986	Paris, France
March 26–28, 1987	Baden/Vienna, Austria
August 23–31, 1987	3rd Congress; Chicago, USA
April 28–29, 1988	Lausanne, Switzerland
August 25–27, 1988	Budapest, Hungary
May 18–19, 1989	Modena, Italy
September 5–8, 1989	4th Congress; Tokyo, Japan
September 21–22, 1990	Gothenburg, Sweden
April 2–5, 1991	Cambridge, UK
August 18–23, 1991	5th Congress; Rome, Italy
February 15–17, 1992	San Francisco, USA
October 29–30, 1992	Lille, France
April 16–18, 1993	Toronto, Canada
September 5–10, 1993	6th Congress; Berlin, Germany
June 5–8, 1994	Amsterdam, The Netherlands
February 17–20, 1995	Cancun, Mexico
October 29–Nov 3, 1995	7th Congress; Sydney, Australia
March 9–10, 1996	New Delhi, India
October 4–5, 1996	Reykjavik, Iceland
April 25–27, 1997	São Paulo, Brazil
August 17–22, 1997	8th Congress; Jerusalem, Israel
May 21–23, 1998	Istanbul, Turkey
September 13–18, 1998	Munich, Germany
April 12–14, 1999	Beijing, China
August 15–20, 1999	9th Congress; Vancouver, Canada
April 4–7, 2000	Newcastle upon Tyne, UK
October 13–15, 2000	Pôrto Alegre, Brazil

Table 142.2 IPA Presidents

M. Bergener, Germany (1982)
G. Bucht, Sweden (1987)
K. Hasegawa, Japan (1989)
S. Finkel, USA (1991)
B. Steen, Sweden (1993)
R. Levy, UK (1995)
B. Reisberg, USA (1997)
E. Chiu, Australia (1999)

The regional meetings and congresses have been the lifeblood of the organization and have brought together diverse professionals who are interested in our field. These include geropsychiatrists, neurologists, geriatric general and family

physicians, psychologists, geropsychiatric nurses, social workers, occupational therapists and others with an interest in our field. Presidents of the organization have until now been drawn from the disciplines of psychiatry and geriatric medicine.

Although these meetings and congresses have been sufficient to serve the growth of our organization, IPA has also created initiatives which have independently served our discipline. One of these initiatives concerns the coveted IPA Research Awards in Psychogeriatrics. The research awards were conceived through the vision of Manfred Bergener and were supported through the generosity of Bayer AG for the first decade, from 1989. These awards, for which submissions are solicited from throughout the world, are given for the best unpublished research submission. Since its inception, the Research Awards Committee has been chaired by Barry Reisberg, with referees from many nations who devote their time, energy and expertise to the exceedingly rigorous review process for these awards.

Winning papers over the course of the years have provided a significant contribution to the body of knowledge and progress in our field. For example, in the first Awards in Tokyo, 1989, subsequently famous research studies on behavioral and psychological symptoms of dementia (BPSD) by Alistair Burns (UK) and on reduction of Alzheimer's disease caregiver stress by Henry Brodaty (Australia) were awarded, in addition to seminal research by Barry Rovner (USA) on agitation in nursing home settings in the USA. Each of these award-winning research entries provided a stimulus for numerous subsequent papers and, more importantly, major changes in the structure of our field. For example, in part as a result of Alistair Burns' work, we now know much more regarding the nature of BPSD and pharmacologic and non-pharmacologic treatments of BPSD. As a result, in part, of Henry Brodaty's award-winning research, Alzheimer's organizations and associated support groups have now proliferated throughout the globe. Barry Rovner's findings led, in part, to changes in the quality of care provided to nursing home residents.

This astoundingly important work, singled out for accolades in IPA's first series of research awards, has resulted in commensurate effects on the careers of these scientists. For example, Alistair Burns is currently the President-elect of IPA and Henry Brodaty is presently the medical director of Alzheimer Disease International. The research awards have remained similarly successful from 1989 to the present, and have clearly not only stimulated the growth of our field but also served to advance the mental health quality of the elderly.

Another modality that IPA has chosen to coalesce growth in our field has been the convening of special meetings. These meetings have typically brought together leading scientists, leading clinicians, government officials, representatives of regulatory agencies and others to coalesce knowledge around a particular subject in our field. For example, a meeting was held in 1994 on "Methodology for drug trials in mild, moderate and severe Alzheimer's disease". This meeting, held in New York, helped to stimulate the approval and understanding of pharmacologic treatments for Alzheimer's disease which have now been approved in many nations around the world.

Another timely special meeting concerned behavioral and psychological symptoms of dementia (BPSD) in 1996. This meeting stimulated worldwide trials of drugs for the treatment

of BPSD and subsequent demonstrations of efficacy. An update special meeting on BPSD was held in 1999. Yet another special meeting, which was held in Geneva in 1996, pulled together current knowledge regarding the diagnosis of Alzheimer's disease from diverse clinical and psychologic, electrophysiologic, neuroimaging, pathologic and biomolecular perspectives. This meeting helped foster the understanding that Alzheimer's disease, like all other major illnesses, is a diagnosis of inclusion as well as of exclusion.

IPA has published a journal, *International Psychogeriatrics*, since 1989, which is an Index Medicus publication and is a leading organ for research and information regarding worldwide activities in psychogeriatrics. Apart from regularly scheduled quarterly issues, *International Psychogeriatrics* has published groundbreaking and syncretic special issues. The special issues have included the proceedings of IPA's special meetings as well as comprehensive issues on other topics, such as a special issue on suicide in the elderly.

Another important publication of IPA is the *Bulletin*, a newsletter which serves as a less formal organ of communication for psychogeriatricians around the world. The IPA also publishes modules, pamphlets and slides on topics of special interest, such as BPSD. The publications further serve IPA's broad educational mission.

Other special and noteworthy activities of IPA include an international visiting junior scholar pilot program, sponsored by Pfizer, which enabled junior psychiatrists and neurologists from less financially endowed research nations to visit wealthier research institutions. For example, physician scholars from Argentina, Brazil, PR China, ROC Taiwan and Russia, were able to work at research centers in the USA, UK and Australia and absorb the most up-to-date methodologies for scientific research.

Yet another important initiative of IPA is its affiliate organization program, which is enabling regional and national organizations to have more ready access to international opportunities and resources and also enable IPA to reach physicians at the "grass roots" level. Equally important in terms of IPA's individual reach is its excellent website (www.ipa-online.org).

Apart from the activities listed in this brief summary, IPA has been involved in numerous other activities and communications in psychogeriatrics. The net result of these activities at the present time was reflected in part by IPA's outstanding Ninth Congress, which was held in Vancouver in 1999. This meeting was the largest meeting ever held in the field of psychogeriatrics. It brought together colleagues from 50 countries that represent IPA's diverse constituency, currently encompassing 75 of the world's nations.

In coming years, IPA seeks to further develop governmental consulting activities in psychogeriatrics, and to further develop its website and affiliate organization program, while maintaining, expanding and improving upon its numerous other activities on behalf of our field.

In summary, through a series of outstanding initiatives and activities, IPA has served the growth of psychogeriatrics worldwide for the past two decades. The goals of these activities continue to be improved mental health as people age, and consequently improved health more generally, throughout the world.