Dementia refers to a syndrome that is characterised by progressive deterioration of cognitive functions. Neuropsychiatric symptoms, such as apathy, agitation, and depression, are also common. With increasing loss of function, a patient is gradually robbed of his or her independence. Eventually, placement in a nursing home may be necessary. Patients with dementia usually survive 7—10 years after onset of symptoms. Dementia places a tremendous burden not only on caregivers, but also on society, and has already been established as one of the major challenges of this century.

Epidemiology refers to the medical science that studies frequencies of disease. Measures of frequency that are often used in epidemiology are prevalence and incidence. The concept of prevalence refers to the number of patients with a disease at a certain moment in time, whereas measures of incidence reflect the number of new cases over time. Although important for health care planners, the knowledge of frequency of disease in itself is not the goal of epidemiology. Rather, the aim is to gain insight into the mechanisms that cause disease, eventually to be able to cure or prevent disease. Therefore, frequencies are studied in relation to determinants, or risk factors. Although marked as “the epidemic of our century”, still surprisingly little is known about the epidemiology of dementia. In this chapter, a brief overview will be given of the epidemiology and risk factors of dementia. Furthermore, we comment on some specific methodological problems associated with studies in dementia.

SYNDROME AND DISEASE

The syndrome of dementia may be caused by various underlying diseases, each characterised by a specific constellation of signs and symptoms in combination with a presumed underlying substrate of neuropathology (fig 1). Alzheimer’s disease (AD) is the most prevalent cause of dementia. It is a neurodegenerative disorder, generally assumed to be caused by neuritic plaques and neurofibrillary tangles accumulating in the brain. The second most prevalent cause of dementia is vascular dementia (VaD), which may be caused by various types of vascular pathology in the brain, such as “large vessel” (large territorial or strategical infarctions) and “small vessel” (lacunes and white matter hyperintensities) disease. Other frequent causes of dementia include frontotemporal lobar degeneration and dementia with Lewy bodies. It is often difficult (if not impossible) to reliably distinguish between subtypes of dementia (we will come back to this subject in the section on methodological issues). Therefore, epidemiological studies often focus on dementia as a whole, sometimes giving separate numbers for the two most important subtypes—AD and VaD.

PREVALENCE

Prevalence is defined as the proportion of a population that has disease at a specific point in time. Prevalence estimates vary highly between studies. These variations may be due to variations in study population—that is, reflect real differences. For example, age is the most important risk factor for dementia. Differences in age between populations will result in different estimates of prevalence. Alternatively, and just as plausible, is the assumption that differences in prevalence estimates are caused by methodological differences, such as study design and diagnostic procedure. One solution to obtain more certainty about the value of prevalence is to combine data from multiple studies in a meta-analysis. Meta-analyses have two advantages. First, small differences due to methodological differences between studies level out. Secondly, and more importantly, the analysis is based on a far larger sample than a single study could ever realise, resulting in more precise estimates.

Pooled estimates of prevalence

In 2000, prevalence data from 11 European population based studies were pooled to obtain stable estimates of prevalence of dementia in the elderly (> 65 years). Age standardised prevalence was 6.4% for dementia (all causes), 4.4% for AD, and 1.6% for VaD. Prevalence of dementia was higher
in women than in men and nearly doubled with every five
year increase in age: 0.8% in the group age 65–69 years and
28.5% at age 90 years and older (fig 2). Of all dementia cases,
54% suffered AD. Prevalence of AD showed the steepest
increase with age, from 0.6% in the group age 65–69 years to
22.2% in the group aged 90 years and older. VaD accounted
for 16% of cases, and prevalence increased with age from
0.3% (65–69 years) to 5.2% (90+ years). More recently,
prevalence rates for dementia were compared among 12
population based European studies. Crude prevalence rates
varied between 5.9% (Italy, the Counselice study) and 9.4%
(the Netherlands, Rotterdam study). Again, an almost
exponential increase with age and a female excess—mostly
after age 75—was described.

**Dementia with young onset**

Most studies on prevalence of dementia focus on subjects
aged over 65 years. Although age is well established as its
most important risk factor, dementia may also affect people
under the age of 65. Few data exist on the prevalence of
dementia in younger people. A recent study in the UK was
designed to determine the prevalence of dementia in people
under the age of 65 in a large catchment area (total
population of 567 500 people) and use these figures to
estimate the number of younger people affected by dementia
in the UK. The prevalence of dementia in those aged 30–64
was 54 per 100 000. For those aged 45–64, the prevalence was
98 per 100 000. Like the studies mentioned above describing
prevalence over the age of 65, there was a strong age
dependency: from the age of 35 years onwards, the
prevalence of dementia approximately doubled with every
class.
than double to well over 16 million patients with dementia. Not only will the number of patients with dementia increase; in the same time span, the working age population will considerably decrease in number (fig 4). While in the year 2000, there was a ratio of 69 working age persons to one demented person, this ratio will decrease to 21:1 in 2050.

In this paragraph, an overview of the prevalence of dementia has been given. Prevalence is determined by both the number of new cases over a given period of time, and by the duration of survival once patients have the disease. Death results in a decrease of prevalence; therefore, diseases that quickly lead to death may have low prevalence, even if they occur frequently, while diseases with long survival have higher prevalence, even if they occur with lower frequency. From the above it follows that studies based on prevalent cases yield associations that reflect the determinants of survival with disease just as much as the causes of disease. This can result in misleading situations—for example, if a new treatment would positively influence the course of dementia by lengthening survival (although not curing the disease), this would result in a higher prevalence. In such a situation, the paradoxical situation may occur that this medication would be positively associated with the prevalence of dementia, and so be misconstrued as a causative agent. For this reason incidence, rather than prevalence, is the desired measure of disease frequency.

INCIDENCE
Incidence refers to the number of new cases over a given period of time. The observed number of new cases depends on the evaluated duration of follow up. To be able to compare studies with varying duration of follow up, incidences per year are usually given. Furthermore, within a given study, length of follow up time may differ between subjects. To profit from all the available information, the length of time at risk is determined for every person. The total length of follow up time is obtained after summing all person-times, and represented as the number of person-years of follow up. Most studies on incidence report incidence rates that are calculated as the number of new cases divided by the person-years at risk. Incidence rates are usually represented as number of new cases per 1000 person-years.

Pooled estimates of incidence
In the same collaborative effort that pooled prevalence data of European studies, data on incidence of dementia of eight population based European studies were compared and pooled. In total, there were 42 996 person-years of follow up with 835 new dementia cases. Of these, 60–70% were diagnosed with AD and 15–20% with VaD. Incidence rates of dementia increased exponentially with age from 2.4 per 1000 person-years in the 65–69 age group, to 70.2 per 1000 person-years in the 90+ age group. Rates among women were higher, especially above the age of 80 (fig 5). The rates continued to increase with age in women, whereas the increase reached a plateau in men at age 85. For AD, findings were comparable, with pooled incidence rates increasing from 1.2 per 1000 person-years among 65–69 year olds to 53.5 among subjects over 90 years old.

Will we be all demented at the age of 140?
The question of whether the incidence rates reach a plateau at a certain age is important, as an exponential increase in
incident AD would suggest that the disease is an inevitable consequence of aging, whereas convergence to a fixed value or a decline could suggest that an element of the population has reduced vulnerability, owing perhaps to genetic or environmental factors. Results with respect to incidence increasing with age have been conflicting, with some studies suggesting an ongoing increase with advancing age, whereas other studies suggest that incidence rates reach a plateau after a certain age. This issue is difficult to resolve, however, as the oldest age groups are always underrepresented, resulting in less precise estimations. The Cache County in Utah, USA, is known for the longevity of its inhabitants. The relatively large proportion of extremely old individuals provides the opportunity to give reliable estimates of incident dementia among the oldest old. There were 185 new cases of dementia (123 AD) among 3308 participants who contributed 10,541 person-years of follow up. The incidence of dementia increased with advancing age from 2 per 1000 person-years in the group aged ≤ 68, to peak with 122 per 1000 person-years in the 90–92 age group, and decline in the 93+ age group (110 per 1000 person-years). The incidence of dementia was higher in females over the age of 80. If incidence rates would indeed plateau at a certain age, then the future public health burden of dementia and AD, albeit still enormous, might be less than previously projected.

**Variation across regions?**

Incidence rates have been found to vary between studies. Methodological issues partly account for these differences, but it is also conceivable that the variable estimates reflect real geographical differences. There are substantial differences in possible risk factors for dementia between regions. Such chronic disease risk factor variation is thought to be responsible for the wide variation seen in other diseases of older age such as cancer and cardiovascular disease (for example, differences between North and South Europe). Given the available evidence for risk factors of dementia and the pronounced variation in vascular risk factors across regions, there could be parallel variation in the incidence of dementia. In fact, the pooled analysis of eight European studies mentioned above suggests a geographical dissociation, with higher incidence rates being found among the oldest old of northwestern countries than among southern countries. To assess variation in incidence within country, the Medical Research Council cognitive function and ageing study (MRC CFAS) compared incidence rates among five sites with different risk patterns and mortality rates. As reported before, incidence was observed to rise with age, particularly above the age of 75, and continued to increase for both males and females into the oldest age groups. However, there was no convincing evidence for variation across sites, and incidence rates did not reflect the variations in the prevalence of possible risk factors in these sites.

**RISK FACTORS**

The estimates of frequency of dementia are important by themselves, as they underline the extent of the health care problem as created by dementia. Although important for health care planners, the frequency of disease in itself is not the most important issue. Rather, we need to gain insight into the mechanisms that cause dementia, to be able to develop therapeutic agents that can slow down or even cure these diseases. Risk factors are studied to find out the basic mechanisms leading to dementia. By influencing these risk factors we hope to be able to modify the course of the disease.

Studies on risk factors for dementia have mainly focused on AD, as it is the most frequent cause of dementia. Age is the most well known risk factor for dementia. Studies of prevalence and incidence of dementia and AD have consistently shown an almost exponential increase with advancing age, in that estimates of both prevalence and incidence double with every five year increase in age. In addition, female sex has repeatedly been shown to be associated with an increased risk of AD, especially at old age. Other risk factors for AD include genetic and vascular factors.

**Genetic risk factors**

Only a small proportion of all individuals with dementia suffers from a familial form of dementia, caused by an autosomal dominant mutation. Mutations in several genes (including Aβ precursor protein, presenilin 1, and presenilin 2) have been shown to cause AD, but these genetic forms of AD account for less than 5% of all cases. The largest proportion of AD cases is therefore “sporadic”. However, genetic factors also seem to influence non-familial cases of AD. The “common disease/common variant” hypothesis postulates that common disorders, such as AD, are also governed by common DNA variants. These variants significantly increase disease risk but are neither necessary nor sufficient to actually cause a specific disorder. Rather, these risk genes display intricate patterns of interaction with each other as well as with non-genetic variables, modifying the risk for a disease. To date, only one such factor has been identified in AD. The apolipoprotein E gene presents in three allelic forms (ε2, ε3, and ε4), of which the ε4 allele is a risk factor for AD. APOE ε4 itself is neither necessary nor sufficient to cause AD, but instead operates as a genetic risk modifier. The well known effect of age on AD is modified by APOE, as age of onset is lower in APOE ε4 positives. In addition, it has been suggested that APOE interacts with vascular risk factors.

**Vascular risk factors**

There is abundant evidence that vascular factors play a role in AD. Vascular risk factors such as hypertension, diabetes mellitus, smoking, and heart disease all have been shown to be associated with AD. Explanations for these associations include: (1) the coincidence of common disorders in the elderly; (2) vascular and cerebrovascular disease precipitating AD; (3) an additive or synergistic (AD + vascular) pathogenesis of dementia; and (4) misclassification of vascular dementia as AD. At this moment, the question about the primary and secondary pathology in AD is unlikely to be answered. The mechanisms linking vascular risk factors to AD remain unclear. Atherosclerosis has been postulated as one common mechanism mediating the association between AD and various vascular risk factors. However, statistical models have failed to demonstrate an important mediating role for atherosclerosis as one common factor. Either the measures of extracranial atherosclerosis are not suitable as proxies for intracranial atherosclerosis, or there are other mechanisms whereby cardiovascular risk factors are associated with AD.

**METHODOLOGICAL ISSUES**

A brief overview has been given of the current knowledge of prevalence, incidence, and risk factors of dementia. Although
progress in understanding dementia is being made, the basic mechanisms causing the majority of dementias are still not known, and satisfying therapeutic options are as yet not available. Studies of dementia are hampered by certain methodological issues inherent to the disorder. These methodological issues may influence the results of studies and be partly responsible for variability in results across studies. Without intending to give a complete overview of the methodological issues associated with the study of dementia, we would like to address briefly four important issues here.

**Diagnostic procedure**

The most important problem with respect to studying dementia and AD is defining the outcome. As yet, there is no single diagnostic test for AD or most of the other types of dementia. The diagnosis of AD is based on clinical criteria, and can be graded as possible, probable, or definite. Several sets of criteria are available, of which the criteria of the National Institute of Neurological and Communicative Disorders and Stroke (NINCDS) and the Alzheimer's Disease and Related Disorders Association (ADRDA), dating from 1984, are the most commonly used. The diagnostic work-up of dementia is time and cost intensive. In large population based studies, it is impossible to assess every subject with a complete diagnostic work-up. Using medical records to identify cases would lead to an underestimation of the number of individuals with dementia, as many of the cases of dementia are never diagnosed in a formal setting.

Therefore, large population based studies usually employ a stepwise approach to identify cases. Most studies use one of two possible stepwise approaches. (1) All subjects are assessed using a screening test. Only those performing below a certain cut off level receive an extensive assessment. A drawback of this approach is the low sensitivity of screening tests. Subjects who are demented but score above cut-off on the screening test are missed. These may include mild cases, and individuals with good cognitive reserve due to, for example, high educational level. (2) A subsample, stratified by certain characteristics such as age, sex, and performance on a screening test, receives an extensive diagnostic assessment. Results are extrapolated to the entire sample. Inherent to this approach is the fact that not all cases will receive an extensive assessment, which may result in lack of precision. The use of different criteria to diagnose dementia, and the variable approaches to operationalise these criteria in large samples, can result in highly varying estimates of frequency. The difficulty of diagnosing mild dementia can lead to an additional problem in incidence studies, as cases that are very mild and therefore not recognised at baseline may be wrongly counted as incident cases at follow up, resulting in biased estimates.

**Insidious onset**

A second—and related—methodological problem inherent to dementia is the insidious onset of the disorder. Neuropathological changes, eventually leading to the clinical syndrome of dementia, may start as early as decades before the disease becomes clinically overt. In analogy with the gradually accumulating neuropathology, the transition from healthy to demented is also gradual, rather than abrupt. The moment when dementia is diagnosed is in fact arbitrary. Moreover, the artificial dichotomy between healthy and demented does not do justice to the continuum of cognitive (dys)function. The concept of mild cognitive impairment (MCI) has been developed to account for the transitional phase between healthy and demented. However, introducing concepts like MCI only shifts the problem, as the borders between healthy and MCI and between MCI and demented remain just as arbitrary and unclear. A possible solution would be to discard the arbitrary distinction between normal and demented, and instead use a continuous outcome, such as a test of cognitive function. This would have several advantages. First, costs and time can be saved as the extensive diagnostic work-up is not necessary anymore. Second, by abolishing the artificial dichotomy into normal and demented, the continuum of cognitive decline is done more justice. This approach also provides the opportunity to study progression of decline within demented individuals.

**Biomarkers**

A third issue reflects the complex relationship between the syndrome of dementia and the underlying diseases. When we talk of AD, we refer to the syndrome that is characterised by progressive memory problems, which usually has an insidious onset, etc. However, at the moment the diagnosis of AD is made, we assume to know the underlying neuropathological substrate—that is, neuritic plaques and neurofibrillary tangles. We assume to know this, because during lifetime, it is impossible to directly measure neuropathology. In fact, post-mortem studies have shown that this assumption in many cases is wrong. In a report of the MRC CFAS of the first 209 subjects (48% demented) who came to necropsy, Alzheimer-type pathology and vascular pathology were equally common, and both correlated with cognitive decline. Most subjects had mixed pathology. Approximately one third of clinically demented patients did not fulfil neuropathological criteria for definite AD, whereas an equally large proportion of non-demented elderly subjects did fulfil these criteria. Neuropathologically, the distinction between different types of dementia, and even between demented and non-demented, seems to be very difficult. The question arises that if it is useful to make clinical distinctions between subtypes of dementia, neuropathology may not even exist. A step towards directly measuring disease, rather than clinical phenotype, would be to take biomarkers as outcome of studies. Both neuroimaging and cerebrospinal fluid can provide useful surrogate markers that give a more direct impression of the pathology. In this way, the possibility of different types of pathology coexisting within one subject is appreciated. For example, magnetic resonance imaging (MRI) measures suggestive of Alzheimer-type pathology and vascular pathology can be evaluated simultaneously.

**Cross sectional versus longitudinal studies**

Studies with a longitudinal design are preferred over studies with a cross sectional design for several reasons. It is conceivable that information about risk factors may be systematically different between patients and controls. Patient data must come from a proxy, who might recall the medical history differently than a proxy of a control or the control himself. In addition, prevalence is determined by both the number of new cases over a given period of time, and by the duration of survival once patients have the disease. In analogy, findings of cross sectional studies can reflect the contribution a risk factor makes to developing dementia as well as to surviving after the dementia starts.
Another important issue in this respect is that risk factors may change over time. The impact of environmental factors, such as smoking, diet, physical activity, and vascular disease, may change over time both within an individual and across birth cohorts. Risk factors such as blood pressure change with ageing. Furthermore, the disease, once it has started, may in turn influence the risk factor. For example, the diet of a demented individual may change, when the person forgets to eat his or her meals on a regular basis. Therefore, the relationship between a risk factor and disease may differ depending on the age the risk factor is measured relative to the outcome.

The age related changes in risk factors make causal inferences as to the development of dementia difficult. Studies of blood pressure in relation to dementia form a good example of how the relationship between risk factors and dementia may be influenced by the moment when the risk factor is measured. There have been conflicting reports, with some studies suggesting that low blood pressure is associated with dementia, whereas others report the opposite, namely that high blood pressure is a risk factor for dementia. Importantly, the age the risk factor is measured relative to the outcome.

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