A diagnosis of dementia is devastating at any age but diagnosis in younger patients presents a particular challenge. The differential diagnosis is broad as late presentation of metabolic disease is common and the burden of inherited dementia is higher in these patients than in patients with late-onset dementia. The presentation of the common degenerative diseases of late life, such as Alzheimer’s disease, can be different when presenting in the fifth or sixth decade. Moreover, many of the young-onset dementias are treatable. The identification of causative genes for many of the inherited degenerative dementias has led to an understanding of the molecular pathology, which is also applicable to later-onset sporadic disease. This understanding offers the potential for future treatments to be tailored to a specific diagnosis of both young-onset and late-onset dementia.

Introduction

Dementia is a major public health concern that is a growing burden owing to an ageing society. However, the high prevalence of dementia in the elderly can overshadow the importance of its occurrence in younger patients. Young-onset dementias can present a substantial diagnostic challenge but can also provide important biological insights that might also be applicable to the more common presentation in older patients. For example, the high prevalence of inherited dementias in younger age-groups has led to the identification of causative genes and subsequent molecular pathology of direct relevance to the more common sporadic disease seen in older patients. The prospect of future treatments targeted at the specific molecular pathological changes of the different dementias makes precise diagnosis essential. In this Review, we discuss the differences between young onset and late onset for the four major dementia diseases: Alzheimer’s disease, vascular disease, frontotemporal lobar degeneration (FTLD), and dementia with Lewy bodies. We also suggest a structured approach to the choice of investigations, building on the “dementia plus” concept; this concept exploits the fact that many of the diseases that cause dementia in young adults also cause additional neurological or systemic features, and the identification of these features can aid diagnosis. A diagnosis of dementia often attracts therapeutic nihilism and so we also include examples of treatable dementias that commonly present to young-onset dementia clinics.

Definitions

The term “presenile dementia”, used widely in the published literature until about 10 years ago, is no longer favoured and the terms “young-onset dementia”, “younger-onset dementia”, and “younger people with dementia” are now commonly used. In this paper, we use the term “young-onset dementia”.

Young-onset dementia is conventionally thought to include patients with onset before 65 years of age. This cutoff point is indicative of a sociological partition in terms of employment and retirement age, but this age has no specific biological significance and there is a range of disease features across this arbitrary divide.

The term dementia as currently defined presents two particular challenges. The first is that standard criteria for dementia require that cognitive impairment is sufficiently severe to compromise social and occupational functioning. The second is that memory must be specifically impaired. A consequence of the first challenge is a delay in a specific diagnosis of the cause of the dementia. This is evident in the diagnostic criteria for Alzheimer’s disease from the National Institute of Neurological and Communicative Disorders and Stroke (NINCDS), which first require that the patient fulfils the criteria for dementia. Thus, before a patient with Alzheimer’s disease can be diagnosed using these criteria, the disease will be well advanced (table 1). Although patients who present with the mild but consistent cognitive decline that accompanies ageing should not be categorised as having dementia, it is increasingly important to make a specific early diagnosis of the cause of cognitive impairment when appropriate, particularly with the possibility of disease-modifying treatments becoming available in the future. The use of the term mild, or minimum, cognitive impairment to describe patients with cognitive impairment that is not of sufficient severity to fulfil criteria for dementia has found widespread support. This term is perhaps most useful when different forms of mild cognitive impairment are recognised (eg, amnestic versus non-amnestic; single versus multi-domain) because these have some value in helping to identify precursor states of specific dementia syndromes. In recently proposed criteria for Alzheimer’s disease, both the importance of early diagnosis and the role of biomarkers irrespective of severity are acknowledged (table 1).

The second challenge, that most criteria for dementia require impairment of episodic memory, has come about because Alzheimer’s disease is both the most common cause of dementia in the elderly and the most studied. The concern here is that patients with progressive cognitive decline without memory impairment (eg, posterior variant Alzheimer’s disease) or with other focal impairments (eg, semantic dementia) might be excluded from the dementia diagnostic algorithm. Thus, the clinical approach should emphasise analysis of the cognitive and neurological syndrome in any patient who...
presents with cognitive decline, and this is particularly the case for young-onset dementia, for which the differential diagnosis is often broad.

Epidemiology

There are few population-based studies on the epidemiology of young-onset dementia and we are not aware of any on cognitive impairment more generally. Harvey and colleagues6 estimated that, in two London boroughs in the UK, the prevalence of dementia with onset between the ages of 30 and 65 years was 54 per 100 000 (95% CI 45–64) and 98 per 100 000 (81–118) between the ages of 45 and 65 years. Alzheimer’s disease was the most common single diagnosis followed by vascular disease and FTLD (figure 1). Using a two-stage postal survey, Ikejima and colleagues7 found a broadly similar prevalence of 42 per 100 000 (39–45) in the Ibaraki prefecture in Japan between the ages of 18–65 years. By contrast with the UK study, the most common cause in the Japanese study was vascular disease, followed by Alzheimer’s disease. There have been several series of convenience samples (eg, groups of patients attending specialist clinics or those in care registries) reported,8–10 which all lend support to the view that Alzheimer’s disease, dementia with Lewy bodies, and vascular cognitive impairment comprise a smaller proportion of cases in younger patients than in the older population. Kelley and colleagues11 analysed a group of patients who were younger than 45 years and found that patients with an onset age of less than 35 years mainly had metabolic causes with substantial overlap with diseases more commonly seen in children and adolescents. By contrast, in individuals who had an onset age of between 35 years and 45 years, most dementias were caused by degenerative diseases.

The extent of overlap between the causes of late-onset and young-onset dementia increases with increasing age, with Alzheimer’s disease, vascular dementia, FTLD, and dementia with Lewy bodies being the most common causes. The younger the onset of dementia, the more likely it is that the patient has a genetic or metabolic disease.12

Table 1: Clinical criteria for dementia and Alzheimer’s disease

<table>
<thead>
<tr>
<th>History</th>
<th>Amnestic MCI</th>
<th>Dementia DSM-IV-TR</th>
<th>Alzheimer’s disease NINCDS-ADRDA</th>
<th>Alzheimer’s disease**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>--</td>
<td>--</td>
<td>40–90 years</td>
<td>--</td>
</tr>
<tr>
<td>Progression</td>
<td>--</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Onset</td>
<td>--</td>
<td>Gradual</td>
<td>Gradual</td>
<td>Gradual</td>
</tr>
<tr>
<td>Memory complaint</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Core: early and significant &gt;6 months</td>
</tr>
<tr>
<td>Activities of daily living</td>
<td>Normal</td>
<td>Impaired</td>
<td>Impaired</td>
<td>--</td>
</tr>
<tr>
<td>Other CNS/general conditions</td>
<td>--</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>Incoordination/gait disturbance</td>
<td>--</td>
<td>--</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>Early seizures</td>
<td>--</td>
<td>--</td>
<td>Absent</td>
<td>Absent</td>
</tr>
</tbody>
</table>

Examination

| Memory impairment | Core | Core | Core | Core: can be isolated |
| General cognitive function | Normal | One or more of: aphasia, apraxia, agnosia, disturbance in executive function | Deficits in at least two areas of cognition | Can be intact or deficits in areas of cognition other than memory |
| Delirium | -- | Absent | Absent | -- |
| General neurological function | -- | -- | Normal | Normal |

Investigation

| MRI | -- | -- | -- | Medial temporal lobe atrophy |
| CSF | -- | -- | -- | Low amyloid, increased tau or phosphorylated tau |
| PET | -- | -- | -- | Bilateral temporoparietal hypometabolism on FDG-PET or abnormal PiB study |
| Genetics | -- | -- | -- | Proven FAD mutation in immediate family |

---not addressed in that set of criteria. Early criteria for Alzheimer’s disease had low specificity and could be used only to diagnose late in the course of the disease. Attempts to refine and accelerate diagnosis included describing a pre-Alzheimer’s disease group: amnestic MCI. However, transition from MCI to Alzheimer’s disease remained poorly defined. Recently published criteria amalgamate MCI and Alzheimer’s disease and are the first to suggest use of biomarkers in diagnosis, reflecting changes in clinical practice.1 MCI=mild cognitive impairment. DSM-IV-TR=Diagnostic and Statistical Manual of Mental Disorders, 4th edition text revision. NINCDS=National Institute of Neurological and Communicative Disorders and Stroke. ADRDA=Alzheimer’s Disease and Related Disorders Association. PiB=Pittsburgh B compound. FDG=fluorodeoxyglucose. FAD=familial Alzheimer’s disease. core=core features. *Probable Alzheimer’s disease=core memory impairment plus one or more supportive investigations.

The major causes of dementia

Alzheimer’s disease, vascular disease, FTLD, and dementia with Lewy bodies are the most common diseases that cause dementia both in the elderly and in
younger patients, although not in those who are younger than 35 years. However, the clinical features of these diseases in younger patients can differ from those seen at a later age.

Alzheimer’s disease

Alzheimer’s first patient was only 51 years at the time of presentation and, for the next 50 years, Alzheimer’s disease was referred to as a presenile dementia. It was the work of Blessed and colleagues that led to the recognition of the importance of the disease in the elderly; these authors showed that the brains of patients who had so-called senile dementia when they died had senile plaques and neurofibrillary tangles that were qualitatively the same as those seen in presenile Alzheimer’s disease. This finding led to the view that the disease was the same regardless of age and the term Alzheimer’s disease has since been used to include all ages. This has been a valuable advance but has masked important differences. One obvious difference is that, compared with elderly individuals, younger patients have fewer comorbidities such as renal disease and heart disease, and lower medication use, which can exacerbate cognitive impairment. Co-existent cerebrovascular disease is also less common in younger patients, whereas there is frequent co-existence of Alzheimer’s disease with vascular disease in the older population.

Autosomal dominant familial Alzheimer’s disease is also more common in individuals with younger onset; sporadic Alzheimer’s disease in individuals younger than 50 years is rare. Although mutations in the amyloid precursor protein (APP) and presenilin-1 and presenilin-2 (PSEN1 and PSEN2) genes, which are associated with familial Alzheimer’s disease, are seen in older patients with Alzheimer’s disease, most patients present below the age of 65 years. In general, individuals with familial Alzheimer’s disease present with features similar to individuals with later-onset sporadic Alzheimer’s disease with prominent episodic memory impairment. This similarity has meant that the insights gained from the studies of pre-manifest and early familial Alzheimer’s disease can be generalised to the more common older-onset sporadic disease. However, unlike patients with sporadic disease, patients with familial Alzheimer’s disease generally have myoclonus, relative preservation of naming, and, in some cases, prominent speech production deficits. Rarely, some patients have features that are not seen in late-onset sporadic disease. In patients with PSEN1 deletions and some point mutations, spastic paraparesis can be a prominent and even a presenting feature years before the onset of cognitive impairment; rarely, a cerebellar ataxia is seen. Mutations in the prion protein (PRNP) gene might cause a clinical syndrome that closely resembles familial Alzheimer’s disease.

There are also important phenotypic variants within the group of patients with younger-onset sporadic Alzheimer’s disease: patients who have non- amnestic deficits collectively comprise about a third of cases who present with young onset compared with about 5% of later-onset presentations. Within the range of non-amnestic disorders, presentations with executive behavioural or language dysfunction are well documented; however, the most frequent phenotype is a biparietal or more posterior biparieto-occipital presentation, so-called posterior cortical atrophy, particularly in those with onset between 50 years and 65 years. The cortical nature of the visual impairment might not be identified, and these patients often have many appointments with opticians and ophthalmologists because of the difficulty they have with locating and perceiving objects. Such patients might not fulfil criteria for Alzheimer’s disease, having relatively preserved episodic memory. The association with the ApoEε4 genotype seen with amnestic Alzheimer’s disease might not be observed in patients with posterior cortical atrophy, hinting at neurobiological differences between these groups of patients.

The ApoEε4 genotype might contribute to a more aggressive clinical disease course in younger patients. A language variant of Alzheimer’s disease, so-called logopenic progressive aphasia, characterised by prolonged word-finding pauses, anomia, and impaired sentence processing, is also more common in younger patients. The extensive anatomical overlap between the logopenic and posterior cortical atrophy syndromes underlines the posterior emphasis of cortical involvement in the younger-onset Alzheimer’s disease variants.

Figure 1: Epidemiology of young-onset dementia

Data from the community study of Harvey and colleagues.
There is an important association between early-onset Alzheimer’s disease and Down’s syndrome. From a neurobiological perspective, recognition of the role of APP over-production due to increased gene dosage with trisomy 21 was an important clue to the identification of the APP gene and the amyloid hypothesis of Alzheimer’s disease pathogenesis. Clinically, people with Down’s syndrome have a substantially increased risk of developing younger-onset dementia after the age of 35 years. Alzheimer’s disease changes at post mortem are essentially universal, while the prevalence of clinical dementia in individuals with Down’s syndrome has been estimated as 15–25% overall and increases steeply with increasing age. Clinical assessment is particularly challenging in this population, particularly as indices of executive and social and emotional functioning might be more important than tests of memory in indicating the onset of clinical dementia.

**Vascular dementia and vascular cognitive impairment**

The term vascular dementia has been problematic for the same reasons as the term dementia, and the term vascular cognitive impairment is preferable. Impairment of episodic memory is less prominent in vascular dementia than in Alzheimer’s disease, particularly in patients with small vessel disease in whom impairment of executive function and cognitive slowing (subcortical dementia) are more common. White matter changes indicative of small vessel disease and lacunar infarcts are commonly seen on MRI scans in elderly individuals and are particularly common in association with Alzheimer’s disease, often indicating “mixed dementia”. In younger patients there is usually, but not invariably, an association with vascular risk factors but intensive investigation might identify rarer causes, including mitochondrial disease or cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL). Amyloid angiopathy is important to recognise as some patients might have an inflammatory component that could be responsive to steroids; lobar microhaemorrhages seen on T2*-weighted MRI might help detection. APP duplications are commonly associated with a prominent amyloid angiopathy with cerebral haemorrhages and seizures. Treatable causes such as cerebral vasculitis are also more commonly found in younger patients than in the elderly.

**Frontotemporal lobar degeneration**

The broad descriptive term frontotemporal lobar degeneration (FTLD) refers to the regional atrophy of a group of non-Alzheimer’s disease degenerative dementias. These dementias are associated with three classic clinical syndromes: behavioural variant FTLD, semantic dementia (a fluent aphasia with loss of word meaning), and progressive non-fluent aphasia (a disorder typified by effortful, non-fluent speech). FTLD accounts for a greater proportion of dementias in younger patients than in elderly patients (figure 1), although this might be partly attributable to ascertainment bias: patients with young-onset dementia are more likely to be studied post mortem and elderly patients with FTLD might be misdiagnosed. This bias might also be associated with the high heritability of FTLD: about 20–40% of FTLD cases are familial in series from specialist referral centres. Among the FTLD syndromes, the behavioural variant is the most heritable and semantic dementia the least heritable, perhaps accounting for recent evidence suggesting that a high proportion of patients with semantic dementia present over the age of 65 years. In cases of autosomally dominantly inherited FTLD in which a mutation has been determined, microtubule-associated protein tau (MAPT) and progranulin (GRN) gene mutations are approximately equally represented. Age at onset tends to be younger in patients with MAPT-associated FTLD than in patients with GRN-associated FTLD, but onset age can be highly variable even within a family. Other rare gene mutations have recently been described (table 2). The extent to which age at onset affects FTLD phenotype is unclear: for the behavioural variant, early-onset and later-onset forms all show a similar range of phenotypes.

FTLD is pathologically heterogeneous and prediction of the underlying pathological process on the basis of the clinical phenotype is generally difficult. Two broad histopathological groupings collectively account for most cases (whether sporadic or inherited): diseases with tau-positive cellular inclusions and diseases with tau-negative, ubiquitin-positive cellular inclusions containing TAR-DNA binding protein (TARDBP; also known as TDP-43). Morphological features of deposition of TARDBP have recently been linked to particular clinical syndromes, suggesting a possible pathogenetic framework that might help resolve the current nosological confusion surrounding FTLD. Among the major FTLD subtypes, the characteristic clinic-anatomical syndrome of semantic dementia has the closest pathological correspondence with tau-negative TARDBP pathology seen in more than 75% of cases; behavioural variant FTLD has wide anatomical and pathological heterogeneity. Non-fluent speech breakdown, particularly in the context of parkinsonism, is more frequently associated with tau pathology, including progressive supranuclear palsy, corticobasal degeneration and Pick’s disease; however, these associations are of limited predictive value in individual patients. The presence of lower motor neuron signs are associated with ubiquitin pathology. A recently identified small subgroup of patients with fused-in-sarcoma (FUS)-positive, TARDBP-negative ubiquitinated inclusions commonly presents with a behavioural syndrome before the age of 40 years; substantial caudate atrophy can be a consistent feature in this group.
Dementia with Lewy bodies and Parkinson’s disease dementia

Dementia with Lewy bodies, the second most common cause of dementia in the elderly, is typically associated with the development of a cognitive syndrome with frontal/parietal involvement, well formed visual hallucinations, and fluctuations, followed by the development of parkinsonism. Dementia with Lewy bodies is characterised pathologically by Lewy bodies, senile plaques, and variable tangle formation. A rare pure form comprises only Lewy bodies with a much younger onset; in a series of nine cases from Japan, eight had an onset before 40 years of age. In other patients, dementia is increasingly recognised as a common feature of advancing Parkinson’s disease, but develops less frequently and with a longer latency in patients with young-onset disease. Patients with early-onset Parkinsonism are more likely to have an underlying genetic cause. Of these, mutations in the parkin (PARK2) gene are not typically associated with dementia, but α-synuclein triplications and mutations in the glucocerebrosidase gene can be associated with prominent cognitive impairment, in some cases resembling classic dementia with Lewy bodies.

Dementia plus syndromes

There are many causes of young-onset dementia, and detailed coverage of them all is beyond the scope of this Review. However, the clinical concept of dementia plus syndromes (i.e., the dementia syndromes in which cognitive impairment is accompanied by additional neurological or systemic features) can be useful to guide a structured approach to clinical diagnosis and investigations. The pattern of cognitive domains that are impaired can be relatively specific in terms of the underlying molecular pathology; for example, posterior cortical atrophy is most frequently associated with Alzheimer’s disease pathology and a verbal semantic

<table>
<thead>
<tr>
<th>Gene</th>
<th>Chromosome</th>
<th>Onset age (years)</th>
<th>Clinical features</th>
<th>MRI features</th>
<th>Pathological changes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alzheimer’s disease</td>
<td>PSEN1</td>
<td>14</td>
<td>Usually &lt;60; typically 35–55</td>
<td>~70% of familial Alzheimer’s disease; resembles sporadic Alzheimer’s disease; can have behavioural presentation; variants associated with spastic paraparesis and ataxia</td>
<td>Disproportionate symmetrical hippocampal atrophy; can have white matter change</td>
</tr>
<tr>
<td></td>
<td>PSEN2</td>
<td>1</td>
<td>Variable, usually 45–65</td>
<td>Very rare (association with Vogt-Geiman ancestry)</td>
<td>Limited information</td>
</tr>
<tr>
<td></td>
<td>APP</td>
<td>21</td>
<td>Usually &lt;65; typically 45–60</td>
<td>~10–15% of familial Alzheimer’s disease; APP duplications recently recognised; associated with early seizes</td>
<td>Disproportionate symmetrical hippocampal atrophy; often prominent white matter change</td>
</tr>
<tr>
<td>Prion disease</td>
<td>PRNP</td>
<td>20</td>
<td>Highly variable</td>
<td>Rarely presents with an amnestic phenotype resembling Alzheimer’s disease; can have rapid course; can be associated with familial fatal insomnia, GSS, or CJD phenotypes</td>
<td>Variable; can have cerebellar atrophy, altered basal ganglia signal on FLAIR</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Deposition of amyloid-containing abnormal prion protein; Alzheimer’s disease-like phenotype associated with prominent vascular involvement</td>
</tr>
<tr>
<td>Behavioural variant FTLD, CBS, PSP</td>
<td>MAPT</td>
<td>17</td>
<td>25–65</td>
<td>5–15% of FTLD; presentation can be cognitive, parkinsonian disorder, or combination; phenotypic variation within families; can have semantic impairment</td>
<td>Frontotemporal atrophy; can have relatively symmetrical anterior mesial temporal lobe atrophy</td>
</tr>
<tr>
<td>Behavioural variant FTLD, PA, CBS</td>
<td>GRN</td>
<td>17</td>
<td>35–90</td>
<td>5–15% of FTLD; wide phenotypic variation within families; often prominent paretal signs; shorter duration of disease than for MAPT mutations</td>
<td>Frontal, temporal, and parietal atrophy on MRI; often strikingly asymmetric</td>
</tr>
<tr>
<td>Behavioural variant FTLD</td>
<td>VCP</td>
<td>9</td>
<td>20–65</td>
<td>Very rare; association with inclusion body myopathy and Paget’s disease, which can precede cognitive complaints; can have sphericentric disturbance, semantic impairment</td>
<td>Frontal or temporal atrophy (can be asymmetric); can have prominent white matter change</td>
</tr>
<tr>
<td>Behavioural variant FTLD</td>
<td>TARDBP (encodes TARDBP)</td>
<td>1</td>
<td>50–75</td>
<td>Very rare (French families); associated with MND; can have semantic impairment</td>
<td>Unclear (frontotemporal hypoperfusion on SPECT)</td>
</tr>
<tr>
<td>Behavioural variant FTLD</td>
<td>Not known</td>
<td>9</td>
<td>40–70</td>
<td>Rare; associated with MND</td>
<td>Frontal atrophy; TARDBP inclinations (MND type) in neurons and glia</td>
</tr>
<tr>
<td>Behavioural variant FTLD</td>
<td>CHMP2B</td>
<td>3</td>
<td>&gt;50</td>
<td>Very rare (Danish family); can have extrapyramidal features, MND</td>
<td>Generalised atrophy; Ubiquitin-positive, TARDBP-negative cytoplasmic inclusions in neurons</td>
</tr>
</tbody>
</table>

Key features of the major familial dementias with autosomal dominant inheritance are summarised. Cognitive dysfunction is generally the hallmark of these diseases at presentation; however, other neurological features can supervene during the course of the disease. CBS=corticobasal syndrome. CJD=Cretzfeldt-Jakob disease. FLAIR=fluid-attenuated inversion recovery sequence. FTLD=frontotemporal lobar degeneration. GSS=Gerstmann-Straussler-Scheinker syndrome. MND=motor neuron disease. PPA=progressive aphasial. PSP=progressive supranuclear palsy. SPECT=single-photon emission computed tomography. APP=amyloid precursor protein. PSEN=presenilin. PRNP=prion protein. MAPT=microtubule-associated protein tau. GRN=granulin. VCP=vasoform-containing protein. TARDBP=TAR-DNA binding protein (also known as TDP-43). CHMP=chromatin-modifying protein. Aβ=amyloid β.

Table 2: Autosomal dominant primary dementia syndromes
memory impairment with ubiquitin-positive, tau-negative inclusions. However, many diseases present with a non-specific memory or fronto-subcortical impairment. In these cases, the additional neurological features or systemic features (dementia plus) can be very informative. Careful examination is mandatory as presence of one of these additional features narrows the differential diagnosis. For example, the presence of a gaze palsy restricts the differential diagnosis and the additional presence of splenomegaly makes a diagnosis of Niemann-Pick disease type C very likely. These dementia plus syndromes are summarised in panels 1 and 2.

Diagnoses not to be missed
A diagnosis of dementia tends to attract therapeutic nihilism and, although treatment is only symptomatic for many patients with degenerative dementias, many other non-degenerative diseases that can present with cognitive impairment or in which dementia is the major or only feature can be successfully treated. The following are examples of diseases that can mimic Alzheimer’s disease and often present in young-onset dementia clinics.

Sleep apnoea
Patients with sleep apnoea can present with memory complaints and, depending on when they are examined, can show cognitive deficits. This presentation is not uncommon: sleep apnoea was reported in 8% of patients presenting to a young-onset dementia clinic. In addition to the typical constellation of snoring, morning headache, and daytime somnolence, more subtle clues can include refractory nocturnal seizures or cerebrovascular disease. The mechanism of cognitive dysfunction might be intermittent hypoaxemia or sleep deprivation. Cognitive improvement, particularly in executive functioning, can be achieved by treating patients with obstructive sleep apnoea, although the effects might not always be substantial.

Transient epileptic amnesia
The syndrome of transient epileptic amnesia is characterised by fluctuations in cognitive function associated with episodes of anterograde amnesia or retrograde amnesia for discrete time periods (eg, vacations or other salient events for which the patient has essentially no recollection). Exacerbation of deficits after sleep is a characteristic feature. Other clinical features of temporal lobe seizures often coexist but these are variable and pure amnestic seizures can occur. Temporal lobe spikes can be evident on a standard electroencephalogram (EEG), but prolonged recording might be needed. There might be evidence of hippocampal damage (altered signal or subtle volume loss) on MRI. Although treatment with anticonvulsants might prevent seizures, and some patients have improvements in cognition, complete resolution of cognitive complaints is unusual.

Limbic encephalitis
The past few years have seen substantial advances in our understanding of limbic encephalitis, a rare syndrome associated with subacute onset of cognitive impairment attributable to medial temporal lobe, amygdala, insula, and orbitofrontal cortical involvement, often accompanied by seizures, psychiatric features, and temporal lobe signal change on MRI. Once infectious causes, including herpes viruses, have been excluded, an immune-mediated

Panel 1: Dementia plus syndromes and associated diseases—neurological features

Ataxia
Spinocerebellar ataxia (particularly types 2, 12, and 17), paraneoplastic diseases, prion diseases (particularly familial forms and variant CJD), DRPLA (common in Japan), fragile x-associated tremor ataxia syndrome, familial British and Danish dementias, mitochondrial disorders, superficial siderosis, neuronal ceroid lipofuscinoses (Kuf’s disease), Niemann-Pick disease type C, multiple system atrophy (dementia usually mild, if present), Alexander’s disease, and multiple sclerosis

Pyramidal signs
Multiple sclerosis, frontotemporal lobar degeneration with motor neuron disease, Alzheimer’s disease (some presenilin mutations), spinocerebellar ataxias, phenylketonuria, familial British and Danish dementias, hereditary spastic paraparesis (SPG4), adrenoleukodystrophy, vanishing white matter disease, polyglucosan body disease, polycystic lipomembranous sclerosing leukoencephalopathy (Nasu-Hakola disease)

Dystonia/chorea
Huntington’s disease (and Huntington’s disease-like syndromes 1–3), Kuf’s disease (characteristic facial dyskinesia), Wilson’s disease, neuroacanthocytosis, pantothenate kinase-associated neurodegeneration (neurodegeneration with brain iron accumulation), Lesch-Nyhan syndrome, DRPLA, corticobasal degeneration, neuroferritinopathy, anti-NMDA receptor-mediated limbic encephalitis, variant CJD

Bucco-lingual mutilation
Neuroacanthocytosis, Lesch-Nyhan syndrome

Akinetic-rigid syndrome
Lewy body disease (dementia with Lewy bodies and Parkinson’s disease dementia), progressive supranuclear palsy, multiple system atrophy (dementia usually mild, if present), Huntington’s disease (particularly juvenile onset), corticobasal degeneration, dementia pugilistica, Wilson’s disease, pantothenate kinase-associated neurodegeneration (neurodegeneration with brain iron accumulation), frontotemporal lobar degeneration with parkinsonism-17, Alzheimer’s disease (usually advanced)

Peripheral neuropathy
Neuroacanthocytosis, cerebrotendinous xanthomatosis, HIV infection, giant axonal neuropathy, alcohol-related diseases, metachromatic leukodystrophy, porphyria, adrenoleukodystrophy, GM2 gangliosidosis, polyglucosan body disease, Krabbe’s disease, sialidosis, Fabry’s disease, mitochondrial disorders, spinocerebellar ataxias (particularly type 3)

Myoclonus or early seizures
Prion disease, Alzheimer’s disease, Lewy body disease, DRPLA, mitochondrial disorders, Gaucher’s disease, GM2 gangliosidosis, neuroserpinopathy, polycystic lipomembranous sclerosing leukoencephalopathy, subacute sclerosing panencephalitis, progressive myoclonic epilepsy syndromes, Kuf’s disease, Lafora body disease, sialidosis

(Continues on next page)
process is most likely. In addition to well established paraneoplastic antibodies (including Hu, Ma2, CV2/CRMP5) targeted against intraneuronal antibodies, a range of antibodies directed against cell-surface antigens (including voltage-gated potassium channels,61 the NMDA receptor,62 the GABA receptor,63 and the AMPA glutamate receptor64) are now associated with a limbic encephalitis phenotype. Certain clinical features can give clues to the causative antibody (eg, older onset with hyponatraemia with voltage-gated potassium channels antibodies; younger female adult/child with dyskinesias in anti-NMDA receptor-mediated limbic encephalitis). In cases of immune-mediated limbic encephalitis, an underlying tumour should be sought and treated if identified; best available evidence suggests that tumour-negative cases should be treated promptly with immunosuppression.65,66 Several antibody-negative patients might also respond to immune modulation and it is likely that further causative antibodies are yet to be determined; Hashimoto’s encephalopathy might fall into this category, with the involvement of thyroid antibodies as an epiphenomenon.65

Clinical assessment

The clinical assessment of a patient with cognitive impairment should be the same regardless of age but the breadth of the differential diagnosis in younger patients, which includes many rare diseases, demands a structured approach. The first objective is to determine the pattern of cognitive and behavioural deficit. The second objective is to determine the involvement of the nervous system more generally. Finally, the general physical examination should not be overlooked because clues to the cause of cognitive dysfunction might lie outside the nervous system. Clinicians should be aware that a patient might initially be referred to non-cognitive specialist clinics; thus, patients with Kuf’s disease or Wilson’s disease might present to a movement disorders specialist at a stage when the cognitive impairment is mild or even absent. However, all patients can present with cognitive impairment when other features are either less salient or absent; for example, the cognitive presentation of multiple sclerosis.66

Cognitive assessment

Much information can be obtained from the bedside assessment of different domains of cognitive function, although widely used instruments such as the mini-mental state examination tend to focus on memory, language, and literacy skills, at the expense of non-dominant hemisphere skills. Testing of executive function at the bedside is often difficult to interpret, but reduced verbal fluency might be a useful non-specific indicator of cognitive dysfunction.67 Dyspraxia and apperceptive agnosia are suggestive of organic disease, implicating dominant and non-dominant parietal function, respectively.

(Continued from previous page)

Gaze palsy

Niemann Pick disease type C (vertical supranuclear; early downgaze loss), Gaucher’s disease (horizontal supranuclear), progressive supranuclear palsy (vertical supranuclear), mitochondrial disorders, spinocerebellar ataxias (particularly type 2), paraneoplastic disorders, Whipple’s disease

Deafness

Superficial siderosis, mitochondrial disorders, familial Danish dementia, alpha mannosidosis, sialidosis

Dysautonomia

Lewy body disease, multiple system atrophy, prion disease (fatal familial insomnia), porphyria, adrenoleukodystrophy, anti-NMDA receptor-mediated limbic encephalitis

The dementia plus syndromes describe patterns of cognitive impairment (dementia) plus additional neurological or systemic features that aid investigation and diagnosis of the underlying disease process. This list cannot be comprehensive. Note that vascular disease, structural disorders, and (para)neoplastic disease can be associated with a wide range of presentations. DRPLA=dentatorubral-pallidoluysian atrophy. CJD=Creutzfeldt-Jakob disease.

A useful bedside distinction can be made between cortical dementia and subcortical dementia. Cortical dementia is characterised by clear errors in specific domains with relative preservation of cognitive speed, and is exemplified by Alzheimer’s disease and semantic dementia.68 By contrast, subcortical dementia68 is characterised by profound slowing of cognition, with a frontal dysexecutive syndrome and impairment of memory retrieval, and is exemplified by progressive supranuclear palsy. In general, the cognitive pattern of dementia plus syndromes is predominantly subcortical. The association of cognitive slowing and behavioural changes has long been associated with disorders of the basal ganglia but similar syndromes are seen with diseases that affect subcortical white matter pathways or ascending projections linking the brainstem to the cortex. These ascending systems are vulnerable to many metabolic disturbances and to drugs and disturbances to these systems account for the pattern of cognitive dysfunction most widely seen with systemic disease.

Although bedside cognitive examination can be informative, a formal neuropsychological assessment is necessary to characterise the patient’s cognitive syndrome in detail. The neuropsychologist uses tests of graded difficulty with well established, age-related normative data so that cognitive performance can be compared across specific domains. It is important to determine premorbid function as far as possible, by use of details of educational attainment, employment history, and reading skills. However, serial assessment is often necessary to be confident of a decline in cognition: neuropsychometry offers the further advantage of quantification of any interval change in performance, bearing in mind that practice effects lead to an improvement in some scores in healthy individuals (eg, IQ scores). The association of cognitive syndromes with particular degenerative dementias is shown in figure 2.69
Panel 2: Dementia plus syndromes and associated diseases—systemic features

Cataracts
Myotonic dystrophy, cerebrotendinous xanthomatosis, mitochondrial disorders, familial
Danish dementia

Splenomegaly
Niemann-Pick disease type C, Gaucher’s disease

Tendon xanthomas
Cerebrotendinous xanthomatosis

Bone cysts
Polycystic lipomembranous sclerosing leucoencephalopathy

Paget’s disease
Valosin-associated frontotemporal lobar degeneration

Renal impairment
Fabry’s disease, Lesch-Nyhan syndrome, mitochondrial disorders

Hepatic dysfunction
Wilson’s disease, Gaucher’s disease, mitochondrial disorders

Respiratory failure
Frontotemporal lobar degeneration and motor neuron disease, Perry syndrome, mitochondrial disease (eg, POLG), anti-NMDA receptor-mediated limbic encephalitis

Gastrointestinal dysfunction
Coeliac disease, Whipple’s disease, porphyria

Anaemia
Vitamin B12 deficiency, neuroacanthocytosis (McLeod’s syndrome), Wilson’s disease, Gaucher’s disease

Skin lesions
Behcet’s disease, systemic vasculitides and connective tissue disease, Fabry’s disease

Metabolic or infectious crises
Vanishing white matter disease, Alexander’s disease, ornithine transcarbamylase deficiency, alpha mannosidosis, porphyria

Hyponatraemia
VGKC limbic encephalitis

The dementia plus syndromes describe patterns of cognitive impairment (dementia) plus additional neurological or systemic features that aid investigation and diagnosis of the underlying disease process. This list cannot be comprehensive. Note that vascular disease, structural disorders, and (para) neoplastic disease can be associated with a wide range of presentations.

Neurological examination
The additional neurological features of a dementia plus syndrome are frequently mild. Careful clinical examination is therefore essential to narrow the large range of potential underlying pathological changes (panels 1 and 2). The range and speed of pursuit and saccadic eye movements can be useful in refining the differential diagnosis (eg, to identify the supranuclear gaze palsy of progressive supranuclear palsy, Niemann Pick type C disease, and Gaucher’s disease). The presence of a jaw jerk or pout, brisk tendon reflexes, and gait apraxia can indicate an underlying vascular disorder. Fasciculations, which can be the clue to lower motor neuron involvement in some types of FTLD, can be restricted to the deltoids and triceps in the absence of long tract signs. Fine myoclonus of the hands in familial Alzheimer’s disease might emerge only when the patient is relaxed or distracted.

General examination
Although systemic ill health (eg, renal or hepatic dysfunction) can be readily apparent from the history or on routine blood tests, a careful general examination is needed to ensure that important clues are not overlooked. Examination of the fundi and blood pressure are mandatory for the identification of vascular disease and vascular risk factors. Inspection of the skin might reveal stigmata of a vasculitic or connective tissue disorder. Subtle splenomegaly in the absence of hepatomegaly is found in adult-onset Niemann-Pick disease type C, and Achilles tendon xanthomata are found in cerebrotendinous xanthomatisos. Systemic findings might also point to an underlying neoplasm; in particular, the breasts and testes should be examined if a paraneoplastic syndrome is a possibility. The patient’s history could be suggestive of obstructive sleep

Behavioural and psychiatric assessment
The behavioural examination begins during history-taking, with an assessment of the patient’s bearing, their interactions with others, and their spontaneous conversation. This assessment is particularly important in patients with behavioural variant FTLD, who might not have deficits on formal cognitive testing but who might make fatuous remarks, perseverate, or have environmental dependency (eg, spontaneously attempting to write with the examiner’s pen). Patients with loss of emotional reactivity might appear inexplicably aloof or hostile. The patient’s approach to testing might also be informative (impulsive in frontal cortical syndromes, slow in subcortical syndromes). Conversely, patients with Alzheimer’s disease generally have a well preserved social appearance but might appear passive during the interview, turning often to their partner to answer questions (the “head turning” sign). In addition to close observation of the patient, it is important to record a history of behavioural and psychiatric (including major mood or psychotic) symptoms, emphasising the need for a corroborating history from an informant who knows the patient well. Some dementias (commonly dementia with Lewy bodies and occasionally FTLD) might present with prominent delusions or other psychotic features; conversely, the profound apathy of negative symptom schizophrenia might mimic a degenerative frontal lobe syndrome. Some patients who present with a static behavioural syndrome and normal imaging might have non-degenerative FTLD phenocopies. Features of REM sleep behaviour disorder should also be sought in the history as this favours a diagnosis of dementia with Lewy bodies.

The dementia plus syndromes describe patterns of cognitive impairment (dementia) plus additional neurological or systemic features that aid investigation and diagnosis of the underlying disease process. This list cannot be comprehensive. Note that vascular disease, structural disorders, and (para) neoplastic disease can be associated with a wide range of presentations.

POLG=polymerase (DNA directed), gamma. VGKC=voltage-gated potassium channel.
apnoea with loud snoring or daytime sleepiness, which can be supplemented by examining for a crowded oropharynx or large collar size.

**Laboratory investigations**

The investigations commonly undertaken in older patients with dementia also apply to young-onset dementia but the broader differential diagnosis mandates a full investigation. The order of investigation follows the general rule of the simplest test first and the most complex and invasive last, which is the order we have detailed below. The extent of the blood tests will, however, depend on the individual patient. For example, neurogenetics is usually confined to those with a positive family history or additional features such as Paget's disease, which is suggestive of FTLD associated with valosin-containing protein (VCP) mutations (table 2).

All patients with young-onset dementia should have structural neuroimaging and CSF examination as recommended by the American Academy of Neurology and European Federation of Neurological Societies guidelines. Decisions about whether to undertake tissue biopsies are based on the clinical phenotype and usually confined to the dementia plus syndromes. Cerebral biopsy is occasionally needed to diagnose cerebral vasculitis. A flowchart for investigations is shown in figure 3, but, because of the wide differential diagnosis, this is not exhaustive. When a diagnosis cannot be established, observation and repeat investigations are often informative.

**Blood tests**

Routine haematological and biochemical blood tests are more useful for detecting comorbidity than for establishing the underlying cause, although metabolic encephalopathies are more likely to occur in younger patients than in older patients. The choice of tests depends on the background and age of the patient, and testing for syphilis or HIV is more relevant in certain settings and, although not routine, should always be considered. Auto-antibodies, antineuronal antibodies, and antibodies implicated in limbic encephalitis should be screened for in patients with rapid-onset dementias or in patients with signs of systemic disease. White cell enzyme and very long chain fatty acid assays are relevant for detection of various metabolic disorders that present in early adulthood, whereas metabolic disorders can best be determined by direct enzyme assay. Rational use of a neurogenetics service relies on an accurate and complete family history in all patients presenting with young-onset dementia, with the caveat that a family history might not always be apparent because of censoring by premature death, non-paternity, or de-novo mutations.

**Neurogenetics**

Neurogenetics has transformed our ability to make precise diagnoses and has extended our understanding of the phenotype of many diseases (eg, leading to the recognition that spastic paraparesis can be associated with some PSEN1 mutations; table 2). Genotyping is often labour-intensive and expensive, although it is anticipated to become less so with novel technologies.

Moreover, at present, screening is impractical in diseases in which many mutations, some family-specific, might be causative. Thus, for some diseases, it is still preferable to establish the diagnosis on the basis of a metabolic profile (eg, increased copper excretion and low ceruloplasmin in Wilson’s disease), and many metabolic disorders can best be determined by direct enzyme assay.

**Imaging**

Historically, the main role of neuroimaging was to exclude a space-occupying lesion, and a CT scan is usually adequate for that purpose. However, MRI offers
sustantial advantages in enabling assessment of signal change and diagnostic patterns of regional brain atrophy. Signal change, particularly in the white matter, best seen with T2 or fluid-attenuated inversion recovery (FLAIR) acquisitions, is an important clue to underlying inflammatory disorders such as multiple sclerosis, vasculitis, limbic encephalitis, or CADASIL (in which there is a characteristic anterior temporal lobe white matter change). Specific patterns of altered signal on FLAIR or diffusion imaging can suggest a prion disease; diffusion imaging is particularly sensitive and should be included if this diagnosis is suspected. MRI sequences sensitive to iron deposition can provide specific clues to several metabolic and genetic disorders (e.g. neuroferritinopathy and pantothenate kinase-associated neurodegeneration). Specific patterns of atrophy, best seen with a volumetric MRI acquisition, can be invaluable in differential diagnosis, reflecting the characteristic patterns of selective neuronal vulnerability (e.g. bilateral hippocampal atrophy in Alzheimer’s disease; asymmetric antero-inferior temporal lobe atrophy in semantic dementia; figure 4). Longitudinal imaging enables changes over time to be visualised and quantified. Metabolic and molecular imaging have an emerging role in the assessment of patients with young-onset dementia. Although fluorodeoxyglucose PET imaging (or single photon emission computed tomography imaging) can show temporoparietal hypometabolism in Alzheimer’s disease, this rarely adds to the visualisation of hippocampal atrophy on structural MRI. By contrast, evidence of frontal hypometabolism can be useful to identify patients with early FTLD (particularly behavioural variant FTLD) and minimum atrophy. PET imaging with ligands such as Pittsburgh B compound (PiB) can show the presence of amyloid, and is likely to emerge as an important diagnostic adjunct in Alzheimer’s disease.”

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Neurophysiology
EEG has tended to fall out of favour in the assessment of cognitive impairment, but the characteristic EEG changes of periodic complexes in some prion diseases and in subacute sclerosing panencephalitis are valuable. Early slowing or loss of alpha rhythm is a feature of Alzheimer’s disease but there is relative preservation of this alpha rhythm in the FTLDs. The EEG can also be used to detect covert epileptiform changes in amnestic syndromes due to partial seizures. Electromyography and nerve conduction studies can be used to identify neuropathies or myopathy in the dementia plus syndromes and can help to establish anterior horn cell dysfunction in patients with FTLD and motor neuron disease.

CSF
A lumbar puncture is recommended by both the American Academy of Neurology and the European Federation of Neurological Society guidelines on investigation of younger patients with dementia. Examination of the CSF can help with the identification of inflammatory causes such as multiple sclerosis, vasculitides, and infections and, together with PCR, might help to identify specific chronic infections such as subacute sclerosing panencephalitis, human herpesvirus 6, Whipple’s disease, cryptococcosis, and tuberculosis. Increasingly, however, the assay of specific proteins has diagnostic value for particular neurodegenerative diseases. The presence of 14-3-3 protein in the CSF is supportive of a diagnosis of Creutzfeldt-Jakob disease and forms part of diagnostic guidelines for dementia. Creutzfeldt-Jakob disease is also characterised by high concentrations of CSF tau. Decreased amyloid β₁–₄₂ and increased tau concentrations have good sensitivity and specificity for Alzheimer’s disease and are also predictive at the mild cognitive impairment stage, and therefore have been incorporated into recent diagnostic criteria for Alzheimer’s disease.

Tissue biopsies
Tissue biopsy might be needed to establish the diagnosis in a few cases, as directed by the clinical features. Skin biopsy (including apocrine sweat glands) might be used to detect abnormal accumulations in storage diseases and in CADASIL, whereas culture of skin fibroblasts can confirm the diagnosis of Niemann-Pick disease type C. Muscle biopsy including histochemistry and respiratory enzyme analysis can confirm a mitochondrial disorder. Tonsillar biopsy can confirm the diagnosis in patients with suspected variant Creutzfeldt-Jakob disease in the absence of a pulvinar sign on MRI. Brain biopsy, however, might be necessary in exceptional cases of young-onset dementia if there is suspicion of a treatable (inflammatory or infectious) process, the diagnosis cannot be made by other means, or if potentially toxic treatment is contemplated. Unless a focal lesion is present, this is necessarily a “blind” procedure, generally from the non-dominant frontal lobe. A full thickness open biopsy including cortex, white matter, and meninges should be done by a neurosurgical team experienced in the technique. Disposable instruments should be used in cases in
which prion risk is thought to be clinically significant. A specific diagnosis can be anticipated in more than 50% of cases and a treatable process in about 10% of cases,

Conclusions

In this Review, we have focused on the clinical approach to young-onset dementia, emphasising the breadth of the differential diagnosis and the need for a structured approach based on all clinical features. The management of young-onset dementia presents challenges that differ from those of older patients. By definition, all are of employable age; many might be the main earner and will often be a parent of young children. The burden of genetic disease in young-onset dementia is also higher than in late-onset dementia. The general issues of management of challenging behaviours, non-pharmacological strategies, and carer support and education all apply in young-onset dementia. However, within the many diseases that can present as young-onset dementia, many demand specific treatment and some curative treatments, necessitating careful investigation. Furthermore, with the dissection of the molecular pathology of the degenerative dementias, specific diagnosis will be essential as disease-modifying therapies become available. Much of our understanding of the pathogenesis of the degenerative dementias has been driven by the identification of genetic mutations causing early-onset familial disease. Thus, the development of transgenic mouse models for Alzheimer’s disease followed directly from the discovery of mutations in the APP and PSEN genes. The extent to which early-onset familial Alzheimer’s disease is typical of late-onset sporadic disease is an important area of future research.

Contributors

All authors contributed to the writing and reviewing of this paper.

Conflicts of interest

NCF has served on the scientific advisory boards of Alzheimer’s Research Forum, Alzheimer’s Society and Alzheimer’s Research Trust. He holds a patent for QA Box that might accrue revenue. In the past 5 years, his research group has received payment for consultancy or for conducting studies from Abbott Laboratories, Eli Lilly, GE Healthcare, IXICO, Lundbeck, Pfizer Inc, Sanofi-Aventis, and Wyeth Pharmaceuticals. MNR, CJM, JMS, and JDW have no conflicts of interest.

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