Approach to atypical Alzheimer’s disease and case studies of the major subtypes

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Alzheimer’s disease (AD) has long been recognized as a heterogeneous illness, with a common clinical presentation of progressive amnesia and less common “atypical” clinical presentations, including syndromes dominated by visual, aphasic, “frontal,” or apraxic symptoms. Our knowledge of atypical clinical phenotypes of AD comes from clinicopathologic studies, but with the growing use of in vivo molecular biomarkers of amyloid and tau pathology, we are beginning to recognize that these syndromes may not be as rare as once thought. When a clinician is evaluating a patient whose clinical phenotype is dominated by progressive aphasia, complex visual impairment, or other neuropsychiatric symptoms with relative sparing of memory, the differential diagnosis may be broader and a confident diagnosis of an atypical form of AD may require the use of molecular biomarkers. Despite the evolving sophistication in our diagnostic tools, and the acknowledgment of atypical AD syndromes in the 2011 revised diagnostic criteria for AD, the assessment of such patients still poses substantial challenges. We use a case-based approach to review the clinical and imaging phenotypes of a series of patients with typical and atypical AD, and discuss our current approach to their evaluation. One day, we hope that regardless of whether a patient exhibits typical or atypical symptoms of AD pathology, we will be able to identify the condition at a prodromal phase and institute a combination of symptomatic and disease-modifying therapies to support cognitive processes, function, and behavior, and slow or halt progression to dementia.

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Introduction

As our ability to measure biomarkers specific to certain neurodegenerative diseases has advanced, it has become increasingly clear that we need to separate neuropathological disease entities (the “disease pathology”) from clinical syndromes of neuropsychiatric dysfunction (the “illness” or “clinical syndrome”). The neuropathological disease known as Alzheimer’s disease (AD), with hallmark amyloid-β neuritic plaques, tau neurofibrillary tangles, and neuronal loss, is well-known to manifest clinically as a variety of diverse syndromes. The most common clinical syndrome associated with AD pathology is the “typical” amnesia-predominant multidomain dementia syndrome that likely begins in most cases as amnesic mild cognitive impairment (MCI). In fact, this form of the illness is so common that for many years the diagnostic criteria required impairment of memory plus impairment of one or more other domains of cognitive function.1 Clinicopathologic reports have called attention to the heterogeneity of AD,2–4 including “atypical” variants of AD,5–7 such as posterior cortical atrophy (PCA), sometimes known as the “visual variant” of AD,
aphasic variants of AD, a behavioral-compartmental (“frontal”) variant of AD, a dysexecutive variant, and even motor variants, including cases that meet clinical criteria for corticobasal syndrome (CBS). As such, AD might be considered the “great imitator” of our time, at least when it comes to other neurodegenerative diseases.

Clinicopathologic studies provide the foundation for knowledge of atypical clinical phenotypes of AD, but with the growing use of specific in vivo molecular biomarkers of amyloid and tau pathology, we are beginning to recognize that these syndromes may not be as rare as once thought. Approximately one-third of patients with AD and onset of symptoms before age 65 present atypically with primary cognitive dysfunction in a domain other than episodic memory,9,10 a phenomenon less common but still encountered in late-onset AD as well.11 Clinical diagnosis is frequently delayed in cases with atypical presentations, and many questions remain about the pathogenesis, risk factors, natural history, and response to treatments in comparison with typical AD.8

In a patient older than 65 with insidiously progressive amnesia, executive dysfunction, and complex visual impairment who has lost independence in daily function to a degree consistent with dementia (the “typical” AD clinical phenotype), many clinicians would likely be highly confident in their diagnosis of probable AD dementia without using molecular biomarkers. In contrast, when a clinician is evaluating a patient whose clinical phenotype is dominated by progressive aphasia, complex visual impairment, or other neuropsychiatric symptoms with relative sparing of memory, the differential diagnosis may be broader and a confident diagnosis of an atypical form of AD may require the use of molecular biomarkers.12 Despite the evolving sophistication in our diagnostic tools, and the acknowledgment of atypical AD syndromes in the revised diagnosis criteria for AD in 2011,13 the determination that a patient with one of these syndromes likely has an atypical form of AD still poses substantial challenges in clinical and research settings. Here we use a case-based approach to review the clinical and imaging phenotypes of a series of patients with typical and atypical AD. First, however, we briefly discuss our current approach to the evaluation of such patients.

Goals of Evaluation and Nomenclature of Diagnostic Summary

When we evaluate a patient, our first goal is to determine whether the overall characteristics and severity of cognitive-behavioral symptoms are consistent with dementia; mild cognitive impairment; encephalopathy (eg, chronic encephalopathies due to immune-mediated or infectious conditions, hormonal or vitamin deficiencies, substance abuse); a learning or attentional disorder; a mood, psychiatric, or sleep disorder; subjective cognitive impairment; or normal cognition. We make this clinical judgment based on all the information gathered during the assessment (eg, assessment of premorbid level/quality and changes in cognitive abilities, activities of daily living, socio-emotional behavior, comportment, sleep, mood, and other neuropsychiatric and medical context), and attempt to grade it, at a minimum, using a severity scale (eg, Clinical Dementia Rating Scale) and a basic cognitive assessment instrument (eg, Montreal Cognitive Assessment score). We will often refer patients at this stage for detailed neuropsychological assessment. This is critical for providing tailored psycho-education and recommendations regarding adaptive planning, safety, and care coordination. Next, we describe the clinical phenotype, including major cognitive, behavioral, and sensorimotor symptoms, and attempt to match it to contemporary syndromic diagnostic criteria. We then consider all of the aforementioned information in order to gauge primary suspected etiology. Finally, we integrate all of the aforementioned information with other indicated diagnostic studies to exclude potentially mimicking conditions (eg, when indicated, serum testing for vitamin B12 deficiency, thyroid hormone disorder, or Hashimoto’s or autoimmune encephalopathy; MRI for mass lesion or vascular cerebral damage/infarct; cerebrospinal fluid (CSF) for voltage-gated channelopathy or paraneoplastic encephalopathy; EEG for subclinical seizures; sleep study for obstructive sleep apnea; urine toxicity or heavy metal screen) and with any available neurodegenerative or other biomarker data and reconsider primary suspected etiology. This approach to the formulation of neurocognitive cases can be summarized as hierarchically determining (1) the patient’s overall functional status along the MCI-dementia spectrum, followed by (2) a description of the major syndrome, followed by (3) a prediction of the likely neuropathology. The case formulation then guides treatment recommendations. When possible, we go through the exercise of stating our confidence in clinical syndrome and suspected etiology before and after diagnostic biomarker testing for the purposes of evaluating the current clinical diagnostic criteria and assessing the utility of current and future diagnostic tests.

A detailed discussion of biomarkers for AD and related neurodegenerative diseases is beyond the scope of this article,14 but we will briefly summarize our current practice. Brain MRI is routinely obtained for most of these patients, or CT with three-dimensional reformating in patients with contraindications to MRI. Regional brain atrophy can provide supportive evidence for the localization of atrophy consistent with neurodegenerative pathology (eg, medial temporal and posterolateral temporoparietal atrophy vs frontal and anterior temporal
Typical Clinical Syndromes Associated with AD

Case 1

Case 1 is a right-handed man who presented at age 62 with a 2-year duration of symptoms. Symptoms included gradually progressive impairment in episodic memory (forgetting important information from recent experiences, including conversations at work and at home, with repetitive asking of questions), in spatial orientation (getting lost in familiar areas), and in judgment and problem solving (no longer able to reason about financial or other decision-making at work or at home), with no reported language, motor, or behavioral-psychiatric symptoms. His impairments resulted in the loss of his job and the need for assistance at home. Medical and family history were unremarkable except for mild hypertension. On exam, the patient demonstrated impaired episodic memory acquisition, retention, and retrieval; impaired complex attention and executive function; and impaired visual construction. Neurological exam was normal. Montreal Cognitive Assessment (MoCA) score was 22; Clinical Dementia Rating (CDR) score was 1 with Sum of Boxes (CDR-sb) of 4.5. Brain MRI scan showed symmetrical atrophy in bilateral rostral hippocampal and medial temporal cortex, medial and lateral parietal cortex, and posterior lateral temporal cortex. At this point, a diagnosis was made of mild dementia, amnesia-predominant syndrome with executive and visual-spatial dysfunction, likely typical AD dementia; the clinician rated his confidence in the clinical syndrome as 100% and the underlying etiology as 95%. An FDG-PET was obtained that showed bilateral inferior parietal, posterior cingulate, and superior temporal hypometabolism. As part of a research study, an amyloid PET scan was visually read as positive. CSF profile of Aβ and tau proteins was highly consistent with underlying AD pathology. These biomarkers brought diagnostic confidence in suspected etiology to 99%. The final clinical diagnosis was dementia, amnesia-predominant multidomain syndrome, highly likely due to AD pathology.

Case 2

Case 2 is a right-handed man who presented at age 64 with 2 years of gradually progressive memory loss. He was no longer able to remember details of conversations with colleagues and now needed to take copious notes. He was having trouble finding his way to places he had been before but to which he traveled infrequently, and needed to rely on prompts from a newly purchased navigation system. There were no reported difficulties with judgment and problem solving, or language, visual, motor, or behavioral-psychiatric function. He was still working as a professor, but the symptoms had resulted in the need for new support systems and greater reliance on an administrative assistant than previously. Medical and family history were unremarkable. On exam, there was normal cognitive test performance except subtle impairment with episodic memory retention and retrieval. Neurological exam was normal. MoCA was 27 (memory); CDR was 0.5, with CDR-sb of 1.5 (memory, spatial orientation, community affairs). Brain MRI scan showed mild left-greater-than-right atrophy in rostral hippocampal and medial temporal cortex with otherwise preserved brain structure (Figure 1). Neuropsychological testing demonstrated impaired verbal and visual memory storage and retrieval (<1 percentile) with below average
acquisition (10 percentile), while other cognitive domains were above average. At this point, a diagnosis was made of MCI, single-domain amnesic subtype. The suspected etiology was AD. The clinician rated his confidence in this syndrome as 100%, and confidence in the etiology as 60%. Additional clinical workup included FDG-PET, which showed left-greater-than-right inferior parietal and superior temporal hypometabolism without obvious posterior cingulate hypometabolism. The clinician then rated his confidence in suspected AD etiology as 80%. Because he and the patient and spouse desired greater confidence, CSF was obtained, which demonstrated a profile of Aβ and tau proteins highly consistent with underlying AD pathology. These biomarkers brought diagnostic confidence to 99%. The final clinical diagnosis was MCI, amnesic syndrome, highly likely due to AD pathology.

This patient exhibits the prototypical prodromal stage of AD, in which the amnesia typical of AD is present. Yet the patient has developed compensatory strategies and is managing to function independently at work and in usual daily activities; thus, he would not be considered to have dementia.13 This is the clinical construct of MCI, originally described in 1999 and subsequently revised to specify cognitive subtypes—amnesic vs non-amnesic.21,22 When a patient experiences gradually progressive amnesia with characteristics suggestive of a “memory storage” problem (as opposed to acquisition or retrieval), there is a strong possibility that the underlying etiology is AD,23 although other neurodegenerative diseases or cerebrovascular disease may also present this way.24 While recent diagnostic criteria incorporating biomarkers into the formulation of likely etiology in patients with MCI specify that these are meant to be research criteria,25,26 we and others are increasingly using them in specialty clinical practice. Using contemporary diagnostic criteria, the patient described here would be classified as having likely prodromal AD25 or MCI due to AD with high likelihood26 or mild neurocognitive disorder likely due to AD.

Atypical Clinical Syndromes Associated with AD

Case 3

Case 3 is a right-handed woman who presented at age 67 with a four-year history of progressive visuospatial impairment. She and her spouse reported difficulty with spatial orientation, including positioning the car correctly in parking spots or in the garage; difficulty with depth perception, including making mistakes on stairs, escalators, revolving doors, and curbs; trouble seeing objects that were “right in front of her,” especially when in the full refrigerator or on a crowded countertop. Her memory was intact. There were no reported symptoms involving judgment and problem solving, language, motor, or behavioral-psychiatric symptoms, except that she was mildly anxious. She was still working as an in-home visiting nurse with some difficulty due to the visual symptoms, but was otherwise functioning independently, doing a variety of community and home activities as usual. Medical and family history were unremarkable. On exam, the only abnormality was visuospatial deficits in figure-copying, clock-drawing, and complex visual perception (difficulty perceiving line drawings of overlapping objects). Neurological exam was unremarkable except for mild oculomotor apraxia, simultanagnosia but no optic ataxia, and very mild left limb apraxia; there was no extrapyramidal dysfunction. MoCA was 26 (visuospatial); CDR was 0.5, with CDR-sb of 1 (spatial orientation, community affairs). The patient had seen an ophthalmologist and was told that there was an inconsistent left partial hemianopia but otherwise normal basic vision. Neuropsychological testing confirmed that, despite normal acuity, complex visual function was significantly impaired (<1 percentile), while other cognitive domains were within normal limits. Brain MRI demonstrated right > left lateral parietal lobe atrophy with preserved medial and lateral temporal lobe structure (Figure 2). FDG-PET showed right > left posterior temporal, parietal, and occipital hypometabolism. At this point, a diagnosis was made of MCI, non-amnestic single domain visual impairment, consistent with posterior cortical atrophy (PCA). Although the clinician was > 90% confident in the clinical syndromic diagnosis, confidence in the likely underlying pathology being AD was 65%. Therefore, CSF was obtained, which showed a profile of Aβ and tau proteins highly consistent with underlying AD pathology. As part of a research protocol, an amyloid PET scan was obtained and visually read as positive. These biomarkers brought diagnostic confidence to 99%
The final clinical diagnosis was MCI, PCA syndrome, highly likely due to AD pathology.

The original description of the posterior cortical atrophy syndrome is usually attributed to D. F. Benson, but multiple earlier case reports describe patients with AD pathology who had prominent early visual disturbances, such as Balint’s syndrome with atypical occipitoparietal pathology. In 1993, a detailed clinical-pathologic report describing “the visual variant of AD” called attention to the severe early visual and spatial impairment with occipito-temporoparietal plaque and tangle neuropathology. Contemporary clinical diagnostic criteria emphasize the presence of progressive visual impairment with relative sparing of memory, language, behavior, and insight; an international work group is currently refining clinical diagnostic criteria. Although contemporary literature largely equates PCA with the visual variant of AD, there are hardly any clinicopathologic studies of PCA with more than 5 cases, with studies suggesting that AD neuropathology may account for 65%, 77%, or even 100%. PCA may also be caused by corticobasal degeneration, Lewy body disease, or, rarely, other neurodegenerative diseases.

**Case 4**

Case 4 is a right-handed woman who presented at age 65 with a 2-year history of progressive language difficulties. She and her daughter described gradually progressive difficulty finding words in conversation, increasing mispronunciation of words, and new difficulty spelling. Her memory was intact. There were no reported symptoms involving spatial or temporal orientation, judgment and problem solving, motor, or behavioral-psychiatric symptoms, except that she reported feeling mildly depressed. She had retired at age 60 but was actively volunteering for 20 hours each week at her local library with little difficulty, and was otherwise functioning independently, living by herself. Medical and family history were unremarkable. On exam, her speech was articulate and fluent at times but with word retrieval difficulties that would reduce fluency along with phonemic paraphasias; she was able to repeat short but not long phrases. Grammar and single word comprehension were normal. The remainder of the office-based cognitive exam was normal except for impairments in spelling, calculation, and verbal list encoding, but retrieval and recognition were normal. Neurological exam was unremarkable except for mild right limb apraxia without rigidity or other extrapyramidal dysfunction. MoCA was 27 (naming, repetition); CDR was 0, with CDR supplemental language box of 0.5. Speech and language pathology assessment demonstrated variable fluency with impairments likely arising during word-retrieval difficulty, anomia, phrase length-dependent repetition impairment, normal verbal grammatical production and comprehension, normal single word comprehension, mildly impaired auditory comprehension for long phrases, normal reading, spelling errors on writing samples but normal grammar, and normal motor speech. Progressive Aphasia Severity Scale (PASS) scores were 0.5 in fluency, 1 in word retrieval, 0.5 in repetition, 0.5 in auditory comprehension, 0.5 in writing; PASS sum of boxes was 3. Neuropsychological testing demonstrated mild verbal encoding impairment (5 percentile), but normal retention and retrieval, with normal visual memory performance and mildly impaired verbal fluency (5 percentile), but normal performance on executive function tasks and tests of other cognitive domains. Brain MRI demonstrated widening of the left Sylvian fissure due to posterior lateral temporal atrophy with preserved medial temporal lobe structure (Figure 3); FDG-PET showed left > right posterior superior temporal and inferior parietal hypometabolism with mild posterior cingulate hypometabolism. A diagnosis was made of MCI, non-amnesic single domain language impairment, consistent with primary progressive aphasia (PPA), logopenic variant (lvPPA). Although the clinician was >90% confident in the clinical syndromic diagnosis, confidence in the likely underlying pathology being AD was 70%. Therefore, CSF was obtained, which showed a profile of Aβ and tau proteins highly consistent with underlying AD pathology. As part of a research protocol, an amyloid PET scan was obtained and visually read as positive. These biomarkers brought diagnostic
confidence to 99% confidence that the underlying disease was likely AD. The final clinical diagnosis was MCI, lvPPA syndrome, highly likely due to AD pathology. Early descriptions of patients presenting with progressive aphasia emphasized the observation that the aphasia sometimes remained isolated for years prior to the development of multidomain impairment and functional loss consistent with dementia. Many of these cases did not show AD pathology, but some did, leading to the idea that this could be an atypical form of AD. Current clinical diagnostic criteria emphasize the presence of progressive language impairment with relative sparing of memory, visual abilities, and behavior. Although many contemporary summaries suggest that the logopenic variant of PPA is essentially equivalent to a language variant of AD, the clinicopathologic investigations of PPA to date indicate that AD neuropathology may only account for about two-thirds of the cases. Moreover, other clinical phenotypes of PPA may be associated with AD pathology.

Case 5
Case 5 is a right-handed woman who presented at age 62 with a 1-year history of progressive cognitive and behavioral symptoms. She reported difficulty with concentration and memory, attributing difficulty at her job as a lab technician to recently diagnosed hypothyroidism. She denied other symptoms. In contrast, her sister reported that her memory was “pretty good,” but that the more notable problems included disorganization and poor judgment and decision-making. She had abruptly left a family gathering for no clear reason, and had recently made several purchases that were beyond her financial capacity (impulsivity). She had developed a new habit of repeatedly checking to make sure her house and car were locked and that she had the keys, and seemed to be collecting pairs of sunglasses (compulsivity). Her sister noticed that she had gained weight and always carried a bag of candy in her purse, a behavior she had never done before (hyperorality). Her sister was concerned that she did not seem to be aware of these unusual behaviors. There were no reported symptoms involving orientation in space or time, language, visual skills, or motor function. She was still working as a lab technician in a hospital but was on probation due to several errors. She was otherwise functioning largely independently, living at home and going on trips with friends, but recently had made 2 errors paying bills, which were out of character and had come to her sister’s attention. Medical and family history were unremarkable except for recently diagnosed hypothyroidism, which was adequately treated. On exam, she had difficulty with performing alternating sequencing and verbal fluency tasks, as well as free recall of words, but was able to correctly retrieve them with cues. Neurological exam was unremarkable except for impersistence when asked to maintain her gaze on an object or hold her arms in the air; there was no extrapyramidal dysfunction. MoCA was 25 (Trails, clock hands, continuous performance test, serial 7s, verbal recall); CDR was 0.5, with CDR-sb of 1.5 (memory, judgment and problem-solving, community affairs); supplemental behavior box score was 1. Social Impairment Rating Scale (SIRS) scores were 0.5 for lack of attention/response to social cues, 0.5 for difficulty with social norms; SIRS sum of boxes was 1. Neuropsychological testing demonstrated borderline performance on tasks of working memory, executive function, verbal and visual recall (5 percentile), but normal encoding and cued recall and recognition. She was impaired on verbal fluency (<1 percentile) but had normal encoding and cued recall and recognition. She was impaired on verbal fluency (<1 percentile) but had normal encoding and cued recall and recognition. She was impaired on verbal fluency (<1 percentile) but had normal encoding and cued recall and recognition. 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study whether clinical features may help to distinguish bvAD from bvFTD. In the end, we believe that clinical features in conjunction with MRI or FDG-PET may improve the probabilistic prediction of AD vs FTLD pathology in a patient with a prominent behavioral syndrome, but molecular biomarkers will likely be necessary to make this discrimination confidently.

Case 6

Case 6 is a right-handed man who presented at age 61 with an 18-month history of gradually progressive movement symptoms followed by cognitive and mood symptoms. He first started having difficulty using his left hand followed by the left leg despite no weakness; sometimes the arm would move “as if it had a mind of its own.” He then developed myoclonic jerks of the left foot and occasionally left arm. He required assistance shaving and getting dressed due to these motor symptoms, and struggled to write and to use utensils and the remote control. Concentration and memory then declined, such that he had to ask people to repeat themselves in conversation and needed reminders for his schedule. He began having difficulty with multitasking. Mild anxiety and depression had developed. He was still working as a manufacturing plant manager with some assistance required and was performing complex activities of daily living largely independently, with the exception of tasks that required the motor functions described above. Medical and family history were unremarkable except for hypercholesterolemia. On examination, difficulties were present with memory encoding, alternating sequences, verbal fluency, and serial 7s. His neurologic exam was notable for mild left-sided extrapyramidal dysfunction with rigidity, right hand dystonia, bilateral ideomotor apraxia, and bilateral agraphesthesia and astereognosia. MoCA was 28 ( Trails, serial 7s); CDR was 0.5, with CDR-sb of 1.5 (memory, judgment and problem-solving, community affairs). Neuropsychological testing demonstrated borderline performance on tasks of working memory, executive function, and verbal and visual encoding (5 percentile), but normal retention and retrieval, and low average performance on verbal fluency (10 percentile), but normal performance in other language domains. Motor speed and dexterity were impaired, more prominently in the left hand (<1 percentile). He also had low average performance (10 percentile) on visual construction tasks. MRI showed right > left precentral and post-central gyrus atrophy (Figure 5). FDG-PET confirmed right > left peri-Rolandic hypometabolism. A diagnosis of MCI, non-amnesic multidomain cognitive impairment with motor impairment consistent with corticobasal syndrome (CBS). The clinician was >85% confident in the clinical syndromic diagnosis, and was...
less than 50% confident in the likely underlying pathology corticobasal degeneration (CBD). Therefore, CSF was obtained, which showed a profile of Aβ and tau proteins highly consistent with underlying AD pathology. As part of a research protocol, an amyloid PET scan was obtained and visually read as positive. These biomarkers brought diagnostic confidence to 99% confidence that the underlying disease was likely AD. The final clinical diagnosis was MCI, non-amnesic multidomain syndrome with predominant motor-cognitive features consistent with CBS, highly likely due to AD pathology.

Although CBS was originally conceptualized as a distinct clinicopathological entity, numerous studies over the past 15 years have highlighted the fact that a substantial minority of cases with classical CBS syndromes arise as a result of AD pathology. Recent studies have shown that a substantial proportion (35%) of patients presenting clinically with CBS ultimately are shown to have AD pathology. Efforts are ongoing to revise clinical diagnostic criteria to better predict pathology, but biomarkers of molecular pathology will almost certainly be an important element of future diagnostic criteria for CBS.

Discussion and Conclusions

The clinical evaluation of AD and other dementias has evolved substantially in the past decade with the advent of a variety of biomarkers of the localization and molecular nature of neurodegenerative diseases. As this has occurred, it has become increasingly clear that we should separate our consideration of the clinical syndrome exhibited by the patient from the suspected underlying pathology, assessing each at least partially independently. Although a progressive amnesic and dysexecutive dementia may be relatively easy to accurately diagnose as likely due to AD pathology, non-amnesic syndromes are less common and present a broader pathological differential diagnosis, and thus are often more difficult to diagnose. In parallel, large pathology investigations have demonstrated that as many as 25% of cases of AD do not conform to the stereotypical progression of neurofibrillary tangle pathology described in the Braak pathology staging scheme. Thus, one of the core principles of behavioral neurology is reinforced: it is not the molecular nature of the lesion that determines the clinical deficit, but rather its localization.

Although fibrillar amyloid plaques are necessary for a pathological diagnosis of AD, the density and distribution of plaques is weakly associated with clinical features in patients with symptoms of the illness. In contrast, detailed neuropathological studies performed more than 2 decades ago showed that the topographical distribution and density of neurofibrillary tangles is closely linked to the clinical phenotype and severity of symptoms. The relationship between tau pathology, regional neurodegeneration, and clinical symptoms has also been reported in atypical forms of AD, including posterior cortical atrophy, behavioral (“frontal”) variant AD, and primary progressive aphasia. The observation that neither the localization of amyloid pathology nor its severity relates to clinical symptoms or markers of neurodegeneration in typical or atypical forms of AD has now been confirmed in vivo using amyloid PET imaging. Amyloid PET studies to date have not demonstrated an ability to distinguish between typical and atypical AD, despite results suggesting higher cortical amyloid burden in apolipoprotein E ε4 non-carriers vs carriers and higher amyloid burden in the parietal cortex in early-onset AD vs late-onset AD. Furthermore, although amyloid PET has revolutionized our approach to the evaluation of patients with suspected AD, like most medical diagnostic tests it may produce false positive or false negative results.

The lack of a PET ligand specific for neurofibrillary tau pathology has rendered it impossible to test the hypotheses regarding relationships of tau to clinical features of AD in vivo. Although CSF tau measures are a validated biomarker of neurofibrillary tau pathology, this biomarker does not enable the localization of tau. As of 2013, this is now changing with the development of new PET ligands to measure tau pathology in vivo.

The first case report of a patient with atypical AD (PCA)
imaged using tau PET demonstrates the co-localization of tau pathology measured in vivo with regional hypometabolism, and the lack of correspondence with regional amyloid.64 This and other recent reports24,75 demonstrate the we are now able to measure the localization and magnitude of both major pathological hallmarks of AD in living patients—a revolution that will almost certainly lead to improved diagnostics and therapeutics.

The question of what factors influence whether a patient may develop typical or atypical AD is largely unanswered. Data from both clinical and pathological studies indicate that younger age is associated with a greater likelihood of an atypical phenotype, as is the absence of an apolipoprotein E ε4 allele.53,59,76 If AD originates and progresses through connections of distributed neural networks in the brain,79-82 the organization of brain networks will shed light on the topographical differences between typical and atypical AD pathology.8 However, it is still unclear why and how a critical node of one brain network rather than another becomes selectively vulnerable to AD pathology in the first place.83 Further investigation is necessary to identify other genetic and environmental drivers of phenotypic diversity in AD, and the mechanisms by which age influences the biology and clinical expression of AD. It is also important to acknowledge that increasing age also makes mixed pathologies more common, such as AD with cerebrovascular disease or AD with cortical Lewy body disease; mixed cases present an additional layer of diagnostic challenge.

The treatment of symptoms of AD may in part be targeted toward specific circuits and the symptoms that arise when they fail, but future therapies will hopefully be able to modify the underlying disease proteinopathies. If this is the case, then determining that a patient’s clinical dementia syndrome is likely due to underlying AD will be a critical factor in guiding the therapeutic approach. One day, we hope that regardless of whether a patient exhibits typical or atypical symptoms of AD pathology, we will be able to identify the condition at a prodromal or preclinical phase and institute disease-modifying therapy, in combination with symptomatic treatments, to slow or halt progression to dementia.

Disclosures

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