Patient with frontal-variant syndrome in early-onset Alzheimer's disease

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ABSTRACT

The clinical manifestation of frontal-variant Alzheimer's disease (FvAD) is not typical, and it is difficult yet necessary to differentiate FvAD from frontotemporal dementia (FTD). We describe a patient with early-onset Alzheimer's disease (AD) who presented with an FvAD-like syndrome and apolipoprotein E ε3/ε4 genotype. A brain amyloid imaging procedure, 11C-Pittsburgh compound B positron emission tomography (PET), supported the final diagnosis of AD. Our present case highlights the clinical variability that characterises early-onset AD. A multimodal approach is crucial when assessing rare forms of dementia.

INTRODUCTION

Atypical frontal-variant Alzheimer’s disease (FvAD), where the brunt of Alzheimer’s disease (AD) pathology is found mainly in the frontal lobes, can present with a dementia syndrome more consistent with frontal-variant frontotemporal dementia (fvFTD) than AD. FvAD, which can take on disinhibited or apathetic forms, is a rare condition associated with early-onset AD.1 We describe here a patient with early-onset AD who clinically presented with fvFTD-like syndrome.

CLINICAL REPORT AND METHODS

A 52-year-old, right-handed woman with 19 years of education was first seen in our geriatric psychiatry department because of memory difficulties combined with progressive and prominent personality changes over the previous 3 years. She had previously been employed as a technician. Her medical history included a partial thyroidectomy due to a bilateral thyroid tumour 1 year ago.

Her caregiver described her onset of memory impairment at age 49, which had developed insidiously. One year later, she showed apparent personality changes including irritability,anhedonia and apathy towards her family, accompanied with memory difficulties, calculation errors and speech disorganisation. She went to a local hospital for testing, and her first MRI scan in February of 2012 revealed mild global atrophy of the cerebral cortex and bilateral hippocampi, with frontal lobe predominance (figure 1A at age 50). The clinical examination revealed no apparent neurological deficits. Cognitive impairment with memory difficulties (Mini-Mental State Examination (MMSE)=22, Montreal Cognitive Assessment (MoCA)=20 at age 50) led to a suspicion of AD and, subsequently, therapy with Aricept was initiated. Six months later, the follow-up outpatient medical records showed that her abnormal behaviours had progressed. The patient became amotivated, lacking the initiative to visit her sick mother. She displayed unusual cooking habits, such as overseasoning of food. Around this time, combination therapy with memantine was added to Aricept. One year later, her caregiver complained of aggravated symptoms, including sluggishness, slowed speech, delayed response, labile mood and odd behaviours such as random hiding of undergarments in bags and other places (MMSE=22, MoCA=18 at age 51). Her second MRI, performed in May of 2013, revealed increased global atrophy of the cerebral cortex and bilateral hippocampi when compared with the first MRI (figure 1B at age 51). At 3 years postonset of symptoms, the patient exhibited increasingly apathetic and strange behaviours, errors in simple arithmetic and poor memory to the extent that it affected her daily functioning (MMSE=12, MoCA=12 at age 52). This was reflected on the third MRI which revealed moderate global atrophy of the cerebral cortex with frontal lobe predominance and bilateral hippocampi shrinkage, leading to suspicion of FTD in March of 2014 (figure 1C at age 52). One month later, she came to our department and received routine laboratory testing, which revealed no evidence of comorbid disease. Our Mini-Mental State Examination revealed a total score of 9 out of 30. The patient exhibited severe impairments in orientation (3/10), attention and calculation (1/5), recall (3/6), language (2/8) and visual construction (0/1). On the MoCA, the patient scored 8 out of...
precursor pathogenic genes for early-onset AD. The direct sequencing showed T/C at the locus of rs429358 and rs7412, which identified APOE ε3/ε4 genotype (figure 1F). In addition, we tested three pathogenic genes for early-onset AD including amyloid precursor protein (APP), Presenilin-1 (PS1) and Presenilin-2 (PS2) using the direct sequencing method, and we did not find any mutation in the three genes. The regional homogeneity (ReHo) analysis of blood oxygen level-dependent (BOLD) functional imaging data showed decreased ReHo (blue) in frontal lobes (figure 1E). Due to apparent personality changes and behaviour dysfunction in the early stages of the disease and late-onset AD risk genotype of APOE ε3/ε4, the patient underwent 11C-Pittsburgh compound B (PIB) PET to provide clarification of diagnosis. This revealed elevated PIB binding in diffuse cortical areas, particularly in the bilateral frontal, parietal, occipital, temporal cortices and posterior cingulated gyrus (figure 1D). The 11C-PIB PET, combined with memory dysfunction as the first clinical manifestation and MRI, confirmed the final diagnosis of AD.

**DISCUSSION**

The patient exhibited insidious memory decline as the first clinical manifestation. The initial MRI revealed a mild global cerebral cortex and bilateral hippocampus atrophy with frontal lobe predominance, and APOE ε3/ε4 genotype showed 3–4 times increase in suffering from late-onset AD. These findings are suggestive of a diagnosis of AD; despite this, we did not find the mutation of early-onset AD pathogenic genes including APP, PS1 and PS2. With disease progression, the patient experienced more prominent personality changes in the early stages of the disease. MRI at this stage revealed moderate atrophy of the cerebral cortex and bilateral hippocampi, with frontal lobe predominance. There was a similar result in that ReHo value decreased in the frontal lobes through the analysis of BOLD functional imaging data. Despite combination therapy with a cholinesterase inhibitor (AChEI) and a non-competitive N-methyl-D-aspartate receptor antagonist (Memantine), cognitive function continued to decline rapidly over the course of 3 years. These particular symptoms seemed suggestive of a diagnosis of FTD. Although the patient began to show mild memory decline and errors in calculation in the early stages, she also exhibited prominent deterioration in personality and social etiquette in the first 2 years. Symptoms indicative of frontal lobe pathology included abulia, apathy, impaired verbal fluency and execution dysfunction. We also considered Lewy body dementia (LBD), given the patient’s abnormal emotional behaviours, but excluded this possibility due to the lack of a Parkinson-like syndrome and definitive hallucinations. Another possibility was a form of rapidly progressive dementia such as prion protein disease. However, the patient was without deficits upon neurological examination or characteristic abnormalities (‘cortical ribbon sign’) on MRI. Beta amyloid is a common biomarker of AD. PIB can specifically bind to extracellular and intracellular fibrous Aβ, allowing PIB PET to be used as an Aβ-specific molecular probe to assess the deposition of Aβ plaques in different brain regions. It can also be used as a non-invasive test method to help dynamic monitoring of protein deposition, playing an important role in early diagnosis of patients with AD. Finally, the patient’s 11C-PIB PET result confirmed the diagnosis of early-onset AD, given her memory decline as the first clinical manifestation, obvious cortex and hippocampus atrophy on MRI, diffuse cortical dysfunction supported by neuropsychological examinations during advanced stages of...
the disease and increased PIB binding in diffuse cortical areas.

FTD is a neurodegenerative disease that affects the frontal and/or temporal lobes, typically affecting individuals between the ages of 45 and 65 years.

Frontal-variant FTD (fvFTD) is the most common form of FTD and is characterised by progressive and prominent deterioration in personality, which takes disinhibited or apathetic forms.

Atypical frontal-variant AD (fvAD), in which the brunt of AD pathology including neuritic plaques and neurofibrillary tangles are found mainly in the frontal lobes, can present with a dementia syndrome more consistent with fvFTD than AD. Our patient appeared to match the apathy symptom profile, which seems to be a behavioural marker indicating a more severe and aggressive clinical decline, with faster cognitive, functional and emotional progression.

The following signs may indicate a clinical diagnosis of fvAD in patients presenting with a fvFTD-like syndrome: genetic signs for AD, including positive family history and APOEε4 allele, and impaired episodic memory performance. As far as laboratory testing and imaging, it may be beneficial to test for cerebrospinal fluid biomarkers or use amyloid imaging procedures such as PIB PET, which aid in the differentiation between fvAD and fvFTD. Tun et al. thought apathy syndrome in AD independently increased the risk of mortality. In typical AD, treatment with AChEIs and/or memantine has been shown to positively affect cognitive, behavioural and daily functioning.

In fvAD, among several psychoactive agents proposed to treat apathy, the best results seem to have been obtained with AChEIs. Selective serotonin reuptake inhibitors (SSRIs) are still a controversial therapeutic approach to treat fvAD. In the case of our patient, however, AChEI seemed ineffective in delaying disease progression.

Behavioural and psychological symptoms of dementia (BPSD) are major and common syndromes in demented persons. FTD was characterised by a much younger mean age of onset than AD and other types of dementia in previous observations. Patients with AD often manifest motor behaviour disturbance, aggressiveness, mood disturbances and psychotic symptoms. Activity disturbances were noted in almost all patients (92.3%) with FTD. Although researchers summarised the clinical characteristics of BPSD in patients with AD and FTD, there remains the possibility of misdiagnosis through observational data. Beach et al. assessed the accuracy of the clinical diagnosis of AD based on autopsy results, or the ‘gold standard’, and demonstrated that depending on the permissiveness of clinical and neuropathological criteria, sensitivity ranged from 70.9% to 87.3% and specificity ranged from 44.3% to 70.8%. Some studies have found that between 12% and 23% of patients diagnosed with AD do not have sufficient AD pathology at autopsy to account for the presence of dementia (‘misdiagnosed’).

Improving diagnostic accuracy in clinical settings may help reduce unnecessary treatment as well as increase the administration of appropriate therapy for patients’ conditions. As a very effective biomarker, beta amyloid PET scan can lead to substantial changes to the methodology of providing diagnoses and intended management plans.

CONCLUSION
FvAD is a rare form of AD which can masquerade as an fvFTD-like syndrome. In the reported case, neither clinical examination nor MRI could definitively differentiate fvAD from fvFTD. Ultimately, these tools, in combination with PIB PET, allowed us to make the diagnosis of fvAD. A multimodal approach is therefore crucial when assessing rare forms of dementia.

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