We report a group of patients presenting with a progressive dementia syndrome characterized by predominant dysfunction in core executive functions, relatively young age of onset and positive biomarkers for Alzheimer’s pathophysiology. Atypical frontal, dysexecutive/behavioural variants and early-onset variants of Alzheimer’s disease have been previously reported, but no diagnostic criteria exist for a progressive dysexecutive syndrome. In this retrospective review, we report on 55 participants diagnosed with a clinically defined progressive dysexecutive syndrome with 18F-fluorodeoxyglucose-positron emission tomography and Alzheimer’s disease biomarkers available. Sixty-two per cent of participants were female with a mean of 15.2 years of education. The mean age of reported symptom onset was 53.8 years while the mean age at diagnosis was 57.2 years. Participants and informants commonly referred to initial cognitive symptoms as ‘memory problems’ but upon further inquiry described problems with core executive functions of working memory, cognitive flexibility and cognitive inhibitory control. Multi-domain cognitive impairment was evident in neuropsychological testing with executive dysfunction most consistently affected. The frontal and parietal regions which overlap with working memory networks consistently demonstrated hypometabolism on positron emission tomography. Genetic testing for autosomal dominant genes was negative in all eight participants tested and at least one APOE ε4 allele was present in 14/26 participants tested. EEG was abnormal in 14/17 cases with 13 described as diffuse slowing. Furthermore, CSF or neuroimaging biomarkers were consistent with Alzheimer’s disease pathophysiology, although CSF p-tau was normal in 24% of cases. Fifteen of the executive predominate participants enrolled in research neuroimaging protocols and were compared to amnestic (n = 110), visual (n = 18) and language (n = 7) predominate clinical phenotypes of Alzheimer’s disease. This revealed a consistent pattern of hypometabolism in parieto-frontal brain regions supporting executive functions with relative sparing of the medial temporal lobe (versus amnestic phenotype), occipital (versus visual phenotype) and left temporal (versus language phenotype). We propose that this progressive dysexecutive syndrome should be recognized as a distinct clinical phenotype disambiguated from behavioural presentations and not linked specifically to the frontal lobe or a particular anatomic substrate without further study. This clinical presentation can be due to Alzheimer’s disease but is likely not specific for any single aetiology. Diagnostic criteria are proposed to facilitate additional research into this understudied clinical presentation.
Introduction

Emerging out of clinical consultations in a tertiary behavioural neurology clinic, we recognized a relatively common pattern among patients presenting with a progressive dementia syndrome characterized by predominant executive dysfunction, early age of onset and positive biomarkers for Alzheimer’s pathophysiology. These patients did not present amnestically like the typical Alzheimer’s disease clinical syndrome. Given the lack of diagnostic criteria and a paucity of published literature describing the characteristics of these patients, they have a prolonged period before diagnosis and are frequently misdiagnosed.

Evidence for a ‘frontal phenotype’ of Alzheimer’s disease has accumulated since the initial reports of a case series (N = 3) of a subgroup of patients with pathologically confirmed Alzheimer’s disease and disproportionate impairment on executive cognitive tests (Trail Making Test A and letter fluency) (Johnson et al., 1999). However, there has been ambiguity in case definitions conflating syndromes (executive versus behavioural) and anatomy (frontal lobe). The International Working Group 2 described a frontal phenotype of Alzheimer’s disease as a progressive behavioural syndrome in the setting of positive Alzheimer’s disease biomarkers that may occur alongside cognitive impairment in executive function (Dubois et al., 2014). In an aim to better represent the clinical and neuroimaging features of such cases, a behavioural/dysexecutive variant of Alzheimer’s disease was then proposed (Ossenkoppele et al., 2015). In this study, Ossenkoppele et al. described that the dysexecutive variant presented with cognitive symptoms 83% of the time and behavioural symptoms 3% of the time, and a behavioural variant presented with cognitive symptoms 53% of the time and behavioural symptoms 25% of the time (12% met criteria for both). The imaging characteristics of these participants included atrophy in the temporoparietal cortex and precuneus, with only subtle frontal lobe atrophy (Ossenkoppele et al., 2015). Whether these two variants represent a clinical spectrum or two distinct variants of Alzheimer’s disease requires further study. This study operationalized a dysexecutive Alzheimer’s disease syndrome using a composite score of executive functions (Digit Span backwards, Trail Making Test Part B, Stroop Color-Word Test and Letter Fluency) being relatively more impaired than a composite score of memory function (Delayed Recall of the Californian Verbal Learning Test or Dutch version of the Rey Auditory Verbal Learning Test and Benson Figure Test or the Visual Association Test) in individuals who had an autopsy and/or biomarker confirmation of Alzheimer’s disease pathology.

In the current study, we aim to provide an operational definition of a progressive dysexecutive syndrome that can be used to describe a clinical presentation to memory clinics that is free from an anatomic description or an underlying aetiological association (e.g. Alzheimer’s disease), as well as free from requirements of particular neuropsychological test results to facilitate further study of this population in a routine clinical context. Neuropsychological testing requirements are not included in the criteria for several reasons: patients may be too impaired at the time of the clinical evaluation to be tested; there exists a lot of variability in which neuropsychological tests are administered, the availability of normative data for younger individuals are lacking as these are often designed to assess cognitive performance in older age individuals; and the fact that most neuropsychological tests require some element of core executive function for performance, resulting in a ‘multi-domain’...
pattern of dysfunction when core executive functions are impaired. This is in line with current criteria in the field that do not specify which cognitive test must be impaired to meet the criteria (McKhann et al., 2011; Dubois et al., 2014).

While it is our experience that many cases presenting with a progressive dysexecutive syndrome eventually are determined to have Alzheimer’s disease as the underlying aetiology, important other causes have been observed in our clinic (e.g. dementia with Lewy bodies, frontotemporal lobar degeneration, Creutzfeldt-Jakob disease and others). However, to raise awareness of the predominant aetiology of the progressive dysexecutive syndrome found in our clinic and to contextualize this syndrome within the broader Alzheimer’s disease literature, we report here only on cases that were on the Alzheimer’s disease continuum based on the new research framework (Jack et al., 2018).

We present an initial description of the clinical presentation, neuropsychological profiles, EEG patterns, genetic results, neuropathologic findings and multimodal neuroimaging characteristics of 55 cases that presented to our clinic with a progressive dysexecutive syndrome that had biomarker or neuropathologic evidence of Alzheimer’s disease. We conclude by discussing these findings to inform aetiologically agnostic diagnostic criteria for a progressive dysexecutive syndrome with added aetiological parameters for Alzheimer’s disease (dysexecutive Alzheimer’s disease) to facilitate additional research into this poorly understood clinical presentation in memory clinics.

### Materials and methods

**Patient consent**

Design and implementation of this single-centre retrospective study met HIPAA guidelines and was approved by the Mayo Clinic Institutional Review Board. Participants signed a research document at their first clinical visit to have their data used in research. Informed consent was obtained for clinical studies (e.g. lumbar puncture and 18F-fluorodeoxyglucose (FDG)–PET) and subsequent research studies (e.g. tau and amyloid PET).

**Participants, clinical features and diagnosis**

A series of participants presenting to an outpatient behavioural neurology clinical practice with a progressive dysexecutive syndrome were collected at Mayo Clinic Rochester between March 2014 and July 2017. A retrospective review of these participants’ clinical Electronic Medical Record was then performed to include those who met the provisional diagnostic criteria for possible or definite dysexecutive Alzheimer’s disease (Box 1) via consensus opinion from four subspecialty trained behavioural neurologists (R.A.T., J.G.-R., H.B. and D.T.J.). This was a modified version of the International Working Group 2 criteria for the frontal phenotype of atypical Alzheimer’s disease (Dubois et al., 2014) and the current NIA-AA research criteria framework (Jack et al., 2018).

One participant did not have *in vivo* biomarkers but had pathology proven Alzheimer’s disease and was also included. Each participant was seen and diagnosed after a visit with a behavioural neurologist. The clinical history and examination were not standardized and were at the discretion of the behavioural neurologist. Behavioural symptoms were probed and documented as positive or negative in the clinical history. Further testing ordered (including formal neuropsychological, EEG and genetic testing) also varied across participants and was based on patient–clinician shared decision-making. Amyloid PET and tau PET were obtained for participants who were also enrolled in our Alzheimer’s Disease Research Center (ADRC).

Exam findings of ideomotor apraxia (e.g. ‘show me how you would hammer a nail’), Luria’s motor series

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**Box 1 Proposed diagnostic criteria**

**Progressive dysexecutive syndrome**

1. A clinical syndrome defined by the presence of persistent, predominant and progressive decline for over 6 months in any core executive cognitive function (i.e. working memory, cognitive flexibility and/or inhibition) in the absence of predominant behavioural features (e.g. would not meet criteria for the clinical syndrome of behavioural variant frontotemporal dementia).

   a. Evidence of impaired executive functions are obtained by patient and/or informant reports in conjunction with formal evaluation of cognitive performance on mentally effortful tasks that require conscious active manipulation of abstract and/or simultaneous information streams.

2. Exclusion criteria include a history of sudden onset or other medical conditions severe enough to account for related symptoms (e.g. primary psychiatric, cerebrovascular, infectious, toxic, inflammatory or metabolic disorders).

**Progressive dysexecutive syndrome due to Alzheimer’s disease (definite dAD)**

- Meets criteria for possible dAD and one of the following:
  1. Increased CSF P-tau
  2. Abnormal tracer retention on amyloid-PET
  3. Alzheimer’s disease autosomal dominant mutation present (PSEN1, PSEN2, APP)
  4. Autopsy consistent with Alzheimer’s disease
(Luria, 1966) and aphasia (using a standard language screen consisting of auditory comprehension, reading comprehension, naming, repetition, narrative picture description and writing) were included in the analysis if a patient’s chart explicitly mentioned the results of these tests. In six cases, participants were referred to a speech pathologist at our institution for detailed speech and language evaluation. Simultanagnosia testing was performed using a combination of Navon figures (Navon, 1977), Ishihara plates (Brazis et al., 1998) and an overlapping figure with five items (Riddoch and Humphreys, 1993). All participants underwent bedside cognitive screening in the form of the Kokmen Short Test of Mental Status (Kokmen et al., 1991).

Neuropsychological testing

Formal neuropsychological testing was performed in 32 out of 55 participants. Neuropsychological testing was not performed in 23 participants due to either travel logistics or severity of cognitive impairment on bedside screening. The neuropsychological battery differed slightly across participants depending on whether they were enrolled as a research participant in the ADRC or undergoing a standard clinical evaluation. The neuropsychological battery for each participant included combinations of the following tests.

Dementia Rating Scale-2 (DRS-2) (Jurica et al., 2001), Rey Auditory Verbal Learning Test (AVLT) (Rey, 1964), Wechsler Memory Scale-Revised (WMS-R) or 3rd Edition (WMS-III) Logical Memory (LM) I and II and Visual Reproductions (VR) I and II subtests (Wechsler, 1987, 1997), Wechsler Adult Intelligence Scale-Revised (WAIS-R) or 3rd Edition (WAIS-III)–Digit Span (DS) and Letter Number Sequencing (LNS) subtests (Wechsler, 1981, 1997), Rey-Osterrieth Complex Figure copy (Osterrieth, 1944), Trail Making Test Part A (Trails A) (Reitan, 1958; Spreen and Strauss, 1998), Trail Making Test B (Spreen and Strauss, 1998), Stroop Test: Word-Reading, Color-Naming and Interference trials (Stroop, 1935), Boston Naming Test (BNT) (Kaplan et al., 1976), Controlled Oral Word Association Test (COWAT) (Ruff et al., 1996) and Category Fluency (Lucas et al., 1998).

Raw scores were converted to age-adjusted standard scores. Mayo Older Americans Normative Studies (MOANS) were used for all tests (Petersen et al., 1992; Ivnik et al., 1997; Lucas et al., 1998; Steinberg et al., 2003; Machulda et al., 2007) with the exception of WMS-III and WAIS-III subtests for which MOANS are not available. For these subtests, published age-adjusted norms from their respective manuals were used. There are three important caveats to these data. First, MOANS norms only extend down to age 56 and four participants were younger than this. In these cases, the youngest MOANS age bracket (56–60) was used to derive normative scores. Second, given the small sample, WAIS-R and WAIS-III scaled scores and WMS-R and WMS-III scaled scores were combined across participants. Finally, two participants had prior neuropsychological evaluations and the data included here reflect their second evaluation (i.e. the evaluations in closest proximity to the imaging and biomarkers). All MOANS and standard scores were converted to z-scores for data presentation.

CSF biomarkers

Lumbar puncture for CSF testing occurred during the week of the initial clinical visit. For Alzheimer’s disease biomarker testing, 2 ml of CSF were collected from each participant at Mayo Clinic, transferred into polypropylene transfer tubes, frozen at −85°C and transported overnight to Athena Diagnostics (Worcester, MA). An enzyme-linked immunosorbent assay employed by Athena Diagnostics was used for CSF analysis for all participants in the present study. An Aβ42: Tau Index <1 is used by Athena and has a sensitivity of 85–94% and a specificity of 83–89% in differentiating clinically diagnosed Alzheimer’s disease from non-Alzheimer’s disease conditions such as vascular dementia and frontotemporal dementia (Hulstaert et al., 1999; Andreasen et al., 2001).

NeuroImaging

FDG–PET images were acquired using a PET/CT scanner (GE Healthcare) operating in 3D mode. Participants were injected in a dimly lit room with FDG, and after a 30-min uptake period, an 8-min FDG scan was performed, which consisted of four 2-min dynamic frames following a low-dose CT transmission scan. Standard acquisition and vendor reconstruction parameters were used. FDG–PET scans were processed using CortexID software (GE Healthcare). The activity in each participant’s PET dataset was normalized to the pons and compared with an age-segmented normative database, yielding z-score 3D-stereotactic surface projection images.

Amyloid-PET imaging was done with Pittsburgh compound B, synthesized on-site with precursor purchased from ABX Biochemical Compounds. Tau PET was carried out with flortaucipir (18F-AV-1451), synthesized on-site with precursor supplied by Avid Radiopharmaceuticals. Image processing methods have been described previously (Jack et al., 2012, 2017). Amyloid and tau PET images were scaled using a cerebellar crus grey matter region of interest (ROI), resulting in standard uptake value ratio (SUVR) images. Previously validated Meta ROIs were used to derive a single value summary measure of amyloid and tau uptake. Positive amyloid PET and tau PET SUVR cutoffs were defined as >1.42 and >1.23, respectively (Jack et al., 2017). Frontal and parietal ROI data were obtained using the Mayo Clinic Adult Lifespan Template (Schwarz et al., 2017). R statistical software was used for exploratory analysis between age of onset, short test of mental status (STMS) and tau PET SUVR.
**Group-wise comparison across Alzheimer’s disease phenotypes enrolled in the Mayo ADRC**

A subset (n = 15) of the participants originally seen in the clinical practice and included in this case series were enrolled in the Mayo Clinic Alzheimer’s Disease Research Center and underwent standardized MRI and FDG–PET imaging. The Mayo Clinic Rochester Alzheimer’s Disease Research Center is a longitudinal cohort study that enrolls subjects from the clinical practice at Mayo Clinic in Rochester, MN. Enrolled participants are adjudicated by a consensus panel consisting of study coordinators, neuropsychologists and behavioural neurologists. These participants were compared to typical amnestic (n = 110), visual (n = 18) and language (n = 7) predominant phenotypes of Alzheimer’s disease dementia enrolled in the Alzheimer’s Disease Research Center that met existing criteria for typical, posterior cortical atrophy or logogenic variant of primary progressive aphasia (Gorno-Tempini et al., 2011; McKhann et al., 2011; Crutch et al., 2017).

Research MR scanning was performed at 3 Tesla and the 3D magnetization-prepared radiofrequency pulses and rapid gradient echo sequences as previously described (Jack et al., 2008). Parameters were: TR/TE/T1, 2300/3/900 ms; flip angle 8°, 26 cm field of view (FOV); 256 x 256 in-plane matrix with a phase FOV of 0.94, and slice thickness of 1.2 mm. These magnetization-prepared radiofrequency pulses and rapid gradient echo parameters have been held invariant since approximately 2008. Hippocampal volume was measured with FreeSurfer (https://surfer.nmr.mgh.harvard.edu/). This structural MRI was used for pre-processing FDG–PET data. Details regarding imaging procedures have been widely reported [see Jack et al. (2017) and references therein].

The FDG–PET image volumes of each subject were coregistered to the subject’s own T1-weighted MRI scan, using a 6 degree-of-freedom affine registration with mutual information cost function. Each MRI scan was then spatially normalized to an older adult template space (Vemuri et al., 2008) using a unified segmentation and normalization algorithm (Ashburner and Friston, 2005) with transforms applied to co-registered FDG–PET images. These spatially normalized images were then intensity normalized to the pons and spatially smoothed with a 6 mm full-width half-maximum Gaussian kernel. These images were then entered into a voxel-wise comparison by Alzheimer’s disease phenotype using SPM12 software (https://www.fil.ion.ucl.ac.uk/spm/) running on MATLAB version R2018a (Mathworks, Natick, MA). Results were considered significant at a voxel-wise FDR corrected P-value of 0.05. Group-wise comparisons of variables were performed using Kruskal–Wallis one-way analysis of variance and pairwise using Wilcoxon rank sum test in R (https://www.R-project.org/) and visualized using the ggplot2 package (Wickham, 2009).

**Genetic testing**

Clinical genetic blood tests included APOE allele status and/or three autosomal dominant genes: Amyloid Precursor Protein (APP), Presenilin-1 (PSEN1) and Presenilin-2 (PSEN2).

**Pathology**

As part of a standardized dissection and sampling protocol (Mirra et al., 1991), formalin-fixed and paraffin-embedded sections of the left hemisphere were taken for immunohistochemical studies. Amyloid plaques and neurofibrillary tangles were immunohistochemically evaluated using antibodies to Aβ (6F/3D; 1:20; Novocastra VectorLabs, Burlingame, CA) and phospho-tau (AT8; 1:1000; Endogen, Woburn, MA) and staged in accordance with recommendations from the National Institute of Aging-Alzheimer’s Association (NIA- AA) and Consortium to Establish a Registry for Alzheimer’s disease (CERAD) guidelines (Mirra et al., 1991; Hyman et al., 2012). Thal amyloid phase was performed using Aβ immunohistochemistry (Thal et al., 2002) and Braak tangle stage was performed using tau immunohistochemistry (Braak and Braak, 1991). Immunohistochemical evaluation of alpha-synuclein (LB509; 1:200; Zymed, San Francisco, CA, USA) did not reveal Lewy body pathology. TDP-43 pathology was evaluated using the MC2085 antibody (rabbit polyclonal, a gift from Leonard Petrucelli, Mayo Clinic) (Zhang et al., 2009). Aging-related tau astrogliopathy was reviewed on tau immunostained sections (Kovacs et al., 2016).

**Data availability**

Data that support the findings in this study are available from the corresponding author upon reasonable request.

**Results**

**Demographics and family history**

Demographics are described in Table 1. There was a slight female predominance of 62%, a mean education of 15.2 years and a mean age of onset of 53.8 years. Thirty out of 55 (55%) participants reported a first or second degree relative with a history of dementia. Only 2 out of 30 relatives reportedly had young onset dementia (<65 years old). For one of the participants with young onset family history, the participant had an APOE ε4 allele present but negative autosomal dominant Alzheimer’s disease genetic testing, and the other participant had no allelic or genetic information available. At least one APOE ε4 allele was present in 14/26 (54%) participants.
Table 1 Broad characterization of progressive dysexecutive syndrome due to Alzheimer’s disease

| Participant demographics (N = 55) |  |
|----------------------------------|---|
| Age of onset (years), mean (range) | 53.8 (48–69) |
| Age at first visit (years), mean (range) | 57.1 (48–71) |
| Education (years), mean (range) | 15.2 (12–20) |
| Gender, No. (%) male | 21/55 (38%) |
| Self-reported family history dementia, N (%) | 30/55 (55%) |
| Young onset family history dementia, N (%) | 2/55 (4%) |

| Genetics | No. with positive test/total tested |
|----------|-----------------------------------|
| APP/PS1/PS2 | 0/8 (0%) |
| APOE ε4 allele carrier status\(^a\) | 14/26 (54%) |

| Clinical and physical exam features | No. with positive finding/total number with documentation of test |
|------------------------------------|---------------------------------------------------------------|
| Parkinsonism | 2/55 (4%) |
| Self-reported RBD | 2/55 (4%) |
| Ideomotor Apraxia | 28/40 (70%) |
| Aphasia | 24/31 (77%) |
| Simultanagnosia | 11/23 (48%) |
| Difficulties with Luria | 39/39 (100%) |

| Neuropsychological testing | Mean (SD) |
|---------------------------|-----------|
| STMS, raw score | 19.9\(^b\) (8.9) |
| Dementia Rating Scale-2, raw, N = 32 | 107.1 (28.9) |
| Dementia Rating Scale-2, z-score, N = 32 | –2.0 (0.9) |
| WAIS-R/III |  |
| Letter Number Sequence, z-score, N = 14 | –1.7 (0.5) |
| Digit Span, z-score, N = 20 | –1.0 (0.7) |
| WMS-R/III |  |
| Logical Memory I, z-score, N = 20 | –2.3 (0.8) |
| Logical Memory II, z-score, N = 20 | –1.9 (0.8) |
| Visual Reproduction I, z-score, N = 17 | –2.4 (0.7) |
| Visual Reproduction II, z-score, N = 17 | –1.8 (0.9) |
| AVLT Trial I, z-score, N = 22 | –1.2 (0.8) |
| AVLT LOT, z-score, N = 22 | –1.4 (0.9) |
| AVLT DR, z-score, N = 22 | –2.0 (0.6) |
| Trail Making Test A, z-score, N = 28\(^c\) | –1.9 (1.1) |
| Trail Making Test B, z-score, N = 22\(^d\) | –2.4 (0.9) |
| Stroop Word, z-score, N = 19 | –1.5 (0.8) |
| Stroop Color, z-score, N = 19 | –1.8 (1.0) |
| Stroop Interference, z-score, N = 19 | –2.1 (0.9) |
| COWAT, z-score, N = 29 | –1.1 (1.2) |
| Category fluency, z-score, N = 22 | –1.6 (1.0) |
| Boston Naming Test, z-score, N = 19 | –0.8 (1.0) |
| Rey-O Figure copy, z-score, N = 24 | –1.8 (1.2) |

| Biomarkers | Median (IQR) |
|------------|--------------|
| Neuron Specific Enolase (pg/ml), N = 35 | 24.4 (21.0–29.0) |
| Aβ42 (pg/ml), N = 51 | 353.7 (279.7–467.4) |
| Total Tau (pg/ml), N = 51 | 522.4 (368.1–996.6) |
| P-Tau181P (pg/ml), N = 51 | 80.1 (60.2–104.7) |
| Amyloid Tau Index, N = 51 | 0.41 (0.25–0.58) |
| Amyloid PET SUVR, N = 22 | 2.41 (2.06–2.33) |
| Tau PET SUVR, N = 20 | 2.30 (1.86–2.77) |
| R Middle Frontal tau PET SUVR, N = 20 | 2.52 (1.64–3.25) |
| L Middle Frontal tau PET SUVR, N = 20 | 2.50 (1.78–3.29) |
| R Superior Parietal tau PET SUVR, N = 20 | 2.52 (1.74–2.94) |
| L Superior Parietal tau PET SUVR, N = 20 | 2.64 (2.06–2.92) |

| EEG | No. with positive finding/total tested (%) |
|-----|------------------------------------------|
| Slowing; epileptiform abnormalities | 14/17 (82%); 4/17 (24%) |

\(^a\) Five participants with homozygous allele, nine with heterozygous allele.

\(^b\) A score of 20/38 on STMS is equivalent to an MoCA of 11/30 or MMSE of 19/30.

\(^c\) Participants who could not complete Trails A (n = 6) were given a MoANS of 1–3.

\(^d\) Participants who could not complete Trails B did not have Trails B attempted and were not included in the z-score. Participants who could not complete Trails B in the maximum allotted time (n = 13) were given a MoANS of 1–3.

AVLT DR = Rey Auditory Verbal Learning Test delayed recall; LOT = Learning Over Trials; RBD = REM Sleep Behaviour Disorder; Rey-O = Rey-Osterrieth Complex Figure; STMS = short test of mental status; WMS-III = Wechsler Memory Scale—3rd edition.
tested. Homozygous APOE ε4 was present in five participants with a mean age of reported onset at 53 years, family history was present in all five (all relatives >65 years old at onset) and one participant had negative genetic testing for APP, PSEN1 and PSEN2. Heterozygous APOE ε4 was present in nine participants with a mean age of reported onset at 52.6 years, a family history in 7/9, young onset family history in one participant and negative autosomal dominant Alzheimer’s disease genetic testing in that individual. No APOE ε4 allele was present in 12 participants with a mean age of reported onset at 52.8 years, a family history in 3/12, no young onset family history and 3 participants with negative autosomal dominant Alzheimer’s disease genetic testing. Overall, there were eight participants with genetic testing for APP, PSEN1 and PSEN2 and all had negative results.

Clinical and physical exam features

All 55 participants presented with cognitive predominant symptoms related to executive functions (example case in Box 2 and Fig. 1). Additionally, behavioural symptoms were present in 16/55 (29%) of participants but were not the predominant symptoms. Fifteen participants had apathy and one participant had impulsive or careless behaviours. Only two participants (3.6%) had mild Parkinsonism on exam, with one of these participants reporting REM sleep behaviour disorder (RBD). Overall, only two participants had RBD. Aphasia was described in 24/31 participants (77%) who exhibited features of logopenic aphasia with word-finding difficulties (circumlocution, lack of specificity and delays for word retrieval), phonological errors or difficulty with repetition. A speech pathology visit confirmed these findings in 6/6 participants. These participants did not meet root criteria for primary progressive aphasia due to language not being the predominant impaired domain or the presenting feature. Moderate to severe difficulties with Luria testing was described in 39/59 participants and 11/23 participants showed signs of simultanagnosia.  

Figure 1 Box 2 Example Case Neuroimaging at presentation (2 years of symptoms) with (A) FDG–PET (Cortex ID, GE Healthcare) showing severe bilateral right > left temporal, parietal, precuneus, posterior cingulate and frontal hypometabolism; (B) MRI Axial T2 Flair and Coronal T1 images with mild right parietal atrophy with no frontal or hippocampal atrophy; (C) 18F-AV1451 tau PET with severe global tau radiotracer uptake; (D) Positive PiB amyloid-PET.  

Box 2 Example case

A 50-year-old factory worker with a past medical history of sleep apnoea presented to the behavioural neurology clinic for a second opinion regarding 2 years of progressive trouble multi-tasking at work leading to job loss. He struggled with coordinating employee shifts, frequently having too many or too few employees scheduled on a given shift. He could no longer count and organize factory inventory. He became extremely anxious when performing more than one activity and had agitated outbursts thought to be related to his frustration with being unable to complete tasks. No other behavioural changes were noted. A neuropsychologic evaluation revealed severe impairment across multiple cognitive domains. He had significant difficulty with letter/number sequencing, block design, perceptual organization, Stroop testing, and he was unable to complete Trails B testing. He was initially diagnosed with bereavement loss (loss of a close friend) and depression due to the discrepancy in his cognitive testing and his relatively preserved functioning in activities of daily living and preserved brain volume.  

His initial neurologic exam revealed cognitive difficulties across multiple domains. He also had difficulty following multistep commands and an inability to perform a motor or written Luria test. Initial FDG–PET and MRI are shown in Fig. 1A and B. His EEG was normal, and his CSF biomarkers were consistent with Alzheimer’s disease (Aβ=42 449, total tau 466 and p-tau 90). He was enrolled in the ADRC and underwent molecular imaging (Fig. 1C and D). Genetic testing was negative for autosomal dominant Alzheimer’s disease genes. Over 2 years his anxiety worsened, and he had difficulty making a sandwich due to trouble organizing what ingredients were needed and in order to add them. On follow-up exam his cognitive testing deteriorated significantly (he was unable to complete testing), and he had prominent acalculia, right/left disorientation, mild paraphasic errors and ideomotor apraxia.

At presentation, nearly all participants were being evaluated for a second or third opinion regarding their diagnosis. In 16/55 participants, the outside working diagnosis was not listed or was not clearly defined (e.g. ‘cognitive impairment of unclear etiology’). In 7/55 participants the outside working diagnosis was fairly correct, labelling it ‘atypical or young onset Alzheimer’s disease’. Misdiagnosis was present in the other 32 participants: 14
diagnosed as a psychiatric (anxiety/depression) disorder, 8 diagnosed as frontotemporal dementia, 4 as autoimmune dementia (no antibodies detected) and failed intravenous steroid trials, 2 as primary progressive aphasia, 1 as corticobasal syndrome, 1 as arsenic poisoning (despite normal lab values), 1 as traumatic brain injury and 1 as chemotherapy-related dementia (lymphoma with no central nervous system involvement).

Neuropsychological testing
Mean z-scores for each neuropsychological test are presented in Table 1. Some participants were administered an abbreviated battery to accommodate the severity of their cognitive deficits (sample sizes provided for each test in Table 1). Nineteen participants discontinued on Trails A (n = 6) or Trails B (n = 13) due to an inability to complete the test in the allowed time (180 and 300 s, respectively). To capture these deficits, we converted these participants’ discontinued scores to a MOANS scaled score of 1 (Machulda et al., 2013), the equivalent of a z-score of −3. The six participants who discontinued on Trails A were not administered Trails B. Multi-domain cognitive impairment was evident, with the magnitude of impairment reflective of the task’s dependence on core executive functions (i.e. working memory, cognitive flexibility and inhibition). Participants had a mean DRS-2 of 107, which is associated with moderate difficulties in basic and instrumental activities of daily living (Fields et al., 2010).

Biomarker confirmation
Biomarker confirmation was established via CSF biomarkers alone (32 participants), CSF biomarkers and positive amyloid and tau PET (15 participants), positive amyloid and tau PET (3 participants), CSF biomarkers and amyloid-PET (2 participants), amyloid-PET (1 participant), autopsy with all biomarker modalities available (1 participant) or autopsy only (1 participant). All participants with CSF testing had an Amyloid Tau Index <1 (median 0.41, IQR 0.25–0.58). Twelve (24%) participants had CSF p-tau below the cut-off value of <61 pg/ml specific to this assay. Five of these participants had tau PET scans, all of which were unequivocally positive with a mean SUVR of 2.05 and are highlighted in Fig. 2. One additional participant with low CSF p-tau and positive tau PET came to autopsy and had pathology proven Alzheimer’s disease.

Electroencephalogram
Seventeen participants underwent a routine 30-min EEG. In 10 participants, the indication for EEG was to work-up early-onset dementia, whereas the 7 remaining participants had atypical episodic symptoms that prompted EEG testing. Of the 14/17 participants with abnormal EEGs, 13 had diffuse slowing (2 participants with superimposed frontal rhythmic delta activity). Three participants had epileptiform discharges arising from the central midline (one participant) and bi-temporal (two participants) regions.

Neuroimaging
All 55 participants underwent MRI and FDG–PET, 22 participants underwent amyloid-PET and 20 participants underwent tau PET (in addition to amyloid PET). Frontal atrophy was minimal on visual inspection, but FDG–PET frontal hypometabolism was prominent in many cases. Lateral parietal, dorsolateral prefrontal, precuneus and posterior cingulate regions were most commonly involved, which visually overlap with regions of 18F-AV-1451 uptake (Figs 2 and 3). To best represent the heterogeneity in the patterns observed, we present neuroimaging findings on the single-subject level in Fig. 3.

Amloid PET was positive for all 22 participants who had imaging and the median SUVR was 2.41 (IQR: 2.06–2.33). The topography followed previously documented Alzheimer’s disease patterns across all participants (Fripp et al., 2008). The 20 participants who underwent Tau PET were all positive and had a median SUVR was 2.30 (IQR: 1.86–2.77). Exploratory analysis reveals that an earlier age of onset was associated with higher global tau PET SUVRs (multiple r² = 0.41; adjusted r² = 0.38, P < .005) and remained significant when controlling for years of symptoms prior to tau PET scan (multiple r² = 0.39, adjusted r² = 0.36, P < .01) (Fig. 4A and B). The middle frontal and superior parietal lobes demonstrated the highest tau PET SUVR loads and were fairly symmetric at the group level (Table 1). Increased global tau PET SUVR was associated with increased cognitive impairment on bedside exam (multiple r² = 0.33; adjusted r² = 0.29, P < .01) (Fig. 4C). Increased global amyloid PET SUVR was not associated with increased cognitive impairment on bedside exam (multiple r² = 0.11; adjusted r² = 0.06, P = .15).

In one participant with longitudinal scans, the severity of 18F-AV-1451 uptake preceded a similar pattern of FDG hypometabolism (Fig. 5).

Neuropathology
Both participants who came to autopsy had a high level of Alzheimer’s disease neuropathologic change per the National Institute on Aging-Alzheimer’s Association 2012 consensus guidelines. The hippocampus was relatively spared compared to frontal and parietal cortex (Fig. 6). The male decedent had evidence of aging-related tau astrogliopathy with clusters of thorn-shaped astrocytes observed in the white matter. Neither participant had evidence of alpha-synuclein-immunoreactive lesions or significant cerebrovascular disease. Microscopic inspection of the amygdala revealed TDP-43 positive neuronal
cytoplasmic inclusions in one of the participants and sub-pial TDP-43 positive neurites in the other participant.

**Group-wise comparison across Alzheimer’s disease phenotypes enrolled in the Mayo ADRC**

Comparing FDG–PET in participants with the executive predominate presentation of Alzheimer’s disease relative to other Alzheimer’s disease phenotypes revealed unique areas of relatively greater hypometabolism in parieto-frontal cortex and relative sparing of the medial temporal lobe (versus amnestic phenotype), occipital (versus visual phenotype) and left temporal (versus language phenotype) (Fig. 7).

A comparison between groups revealed significant differences in age, age at onset, and hippocampal volume (Fig. 8A–C). There was no difference in disease duration, bedside cognition or cortical thickness in Alzheimer’s

**Figure 2 Discordance of CSF p-tau biomarkers and tau PET in five participants who had CSF p-tau in the normal range (< 61 pg/ml). Left: FDG–PET (using Cortex ID, GE Healthcare) with z-score bar at the top. Middle: $^{18}$F-AV1451 tau PET with SUVR bar at the top and individual meta-ROI SUVR below each image. Right: PiB amyloid PET with SUVR bar at the top and individual meta-ROI SUVR below each image. Significant $^{18}$F-AV1451 uptake was present in all five participants (mean SUVR 2.05). CSF p-tau level, tau PET SUVR and mean time lag between CSF and tau PET for each participant can be found in Supplementary Table 1. Bottom participant came to autopsy and was found to have A3, B3, C3 Alzheimer’s disease pathology.**
disease signature regions (Fig. 8D–G). Given the differences in age by phenotype, we repeated the voxel-wise analysis of the FDG–PET images while controlling for age. This altered the statistical thresholds but the anatomic patterns of the relative difference between groups remained the same (data not shown).

**Discussion**

We describe a large group of participants with a progressive dysexecutive syndrome that is attributable to Alzheimer’s disease. In our cohort, nearly all participants were affected during their productive working years at symptom onset, and it took an average of 3 years before they received an accurate diagnosis by a behavioural neurologist. Initial misdiagnosis was common. When clear documentation was available 32 of 39 cases were initially misdiagnosed. Diagnostic confusion can arise from the younger age of onset, the neuropsychological profile, and neuroimaging and biomarker findings. As our example case (Box 2) illustrates, there can exist discordance between a patient’s previous high levels of function with the presence of severe multi-domain impairment on cognitive testing. On average, bedside testing in our cohort was in the moderate to severe dementia range with a mean STMS of 20/38, which is equivalent to a Montreal Cognitive Assessment score of 11/30 (Townley et al., 2019).
and an MMSE score of 19/30 (Tang-Wai et al., 2003). Compared to the degree of clinical impairment, MRI atrophy can be subtle at presentation and most apparent in parietal regions rather than the medial temporal lobe.

This pattern of atrophy has been reported previously (Dickerson et al., 2011; Mendez et al., 2012; Scheltens et al., 2017) but diagnosis can be obscured if one is looking for a typical Alzheimer’s disease pattern. Even when a neurodegenerative etiology is suspected, ancillary tests can create additional diagnostic uncertainty. CSF testing demonstrated normal p-tau in a subset of participants (24% in our cohort), and FDG–PET commonly demonstrated significant frontal hypometabolism (Fig. 7), which can be misidentified as frontotemporal dementia. Based on our provisional diagnostic criteria (Box 1), a diagnosis of dysexecutive Alzheimer’s disease represents a progressive dysexecutive syndrome with a positive biomarker profile along the Alzheimer’s continuum (Jack et al., 2018). According to the NIA-AA research criteria framework, our seven participants with abnormal amyloid (A+) but normal CSF p-tau (T-) and no tau PET available are most accurately described as consistent with Alzheimer’s neuropathologic change rather than definitive progressive dysexecutive syndrome with Alzheimer’s disease. Once tau PET becomes clinically available, it could help address any diagnostic uncertainty in patients with normal CSF p-tau levels (Fig. 2).

Similar to prior studies, atrophy on MRI was mild to moderate in lateral parietal cortices but only mild in frontal cortices (Dickerson et al., 2011; Ossenkoppele et al., 2015).Despite this, FDG–PET and tau PET imaging demonstrated significant hypometabolism and tau radiotracer uptake in the bilateral frontal and lateral parietal cortices. We note that one participant with 3 years of longitudinal data available showed FDG–PET hypometabolism lagging tau PET abnormalities (Fig. 5). Although autopsies are cross-sectional in nature, intra-individual comparisons may provide some insight. Neuropathologic examination revealed significant tau pathology in parieto-frontal cortices with relative hippocampal sparing (Murray et al., 2011) in a pattern mirroring the FDG–PET abnormalities (Fig. 7) and differences in hippocampal volume (Fig. 8C). The interplay and sequencing of amyloid, tau, metabolism, atrophy and clinical symptoms in participants with dysexecutive Alzheimer’s disease are uncertain but are areas of ongoing investigation.

Prior studies have reported conflicting results regarding the involvement of frontal versus parietal regions on molecular imaging in dysexecutive Alzheimer’s disease. One study examined FDG–PET and tau PET in two participants diagnosed with dysexecutive Alzheimer’s disease and showed frontal, parietal and temporal lobe hypometabolism that topographically overlapped with 18F-AV-1451 uptake (Dronse et al., 2017), whereas another study described one participant with dysexecutive/behavioural mixed phenotype with parietal 18F-AV-1451 uptake and

![Figure 4: Age of onset association with tau PET burden](image-url)

**Figure 4: Age of onset association with tau PET burden**

Scatter plots of the 20 participants with tau PET scans: (A) A younger age of onset was associated with a higher tau PET SUVR (multiple $r^2 = 0.41$; adjusted $r^2 = 0.38$, $P < 0.005$). (B) Years of symptoms prior to tau PET was not statistically significant (multiple $r^2 = 0.09$; adjusted $r^2 = 0.04$, $P = 0.21$). (C) Higher tau PET SUVR was associated with lower bedside cognitive testing (multiple $r^2 = 0.33$; adjusted $r^2 = 0.29$, $P < 0.01$).
no frontal uptake (Ossenkoppele et al., 2016). In another, participants with various Alzheimer’s disease phenotypes were described and executive impairment correlated with dorsolateral frontal and lateral parietal tau PET signal (Bejanin et al., 2017). From a molecular imaging standpoint, our study builds upon and reconciles these discordant studies in three ways: (i) although the most common metabolic and tau PET pattern involved both frontal and parietal regions, participants with parietal-predominant involvement also presented with a dysexecutive syndrome, (ii) executive impairment, whether associated with dysexecutive Alzheimer’s disease or other Alzheimer’s disease phenotypes, is associated with lateral prefrontal and parietal tau PET uptake and (iii) like all forms of Alzheimer’s disease, there is significant inter-individual heterogeneity in the pattern of molecular imaging that is associated with a dysexecutive Alzheimer’s disease phenotype (Fig. 3). Future studies should emphasize phenotypic subtypes that may meet the criteria for dysexecutive Alzheimer’s disease.
The factors that drive syndromic diversity and heterogeneity within a specific atypical Alzheimer’s disease syndrome are not well understood. According to the network model of Alzheimer’s disease, genetic and environmental factors may influence regional cortical atrophy patterns (Wolk et al., 2010; Scheltens et al., 2017) and may predispose a selective network to vulnerability (Warren et al., 2012). The pattern of network failure in an individual then drives the clinical and anatomic syndromic diversity (Jones et al., 2017). In our early-onset cohort, the dysexecutive phenotype appeared to be associated with disruption of bilateral parieto-frontal regions, which is consistent with prior early-onset Alzheimer’s disease studies describing executive impairment (Dickerson et al., 2011, 2017; Daianu et al., 2016; Scheltens et al., 2017). We recently described a parieto-frontal tau-PET pattern using a data-driven method and found a similar association with younger age of Alzheimer’s disease onset and tau deposition in this system that spatially overlaps with the working memory network (Jones et al., 2017).

Definitions of executive functions vary, but there is general agreement that there are three core dissociable abilities: (i) working memory (monitoring existing and updating incoming information), (ii) cognitive flexibility (incorporating simultaneous streams of information and set-shifting between mental tasks) and (iii) cognitive inhibitory control (suppressing irrelevant incoming information) (Miyake et al., 2000; Baddeley, 2012; Diamond, 2013). Of note, though simple temporary storage of information is a component of working memory (often referred to as short-term memory), it is the utilization...
and manipulation of this information for completion of tasks/goals that are characteristic of working memory as it relates to executive function (Baddeley, 2012).

Many participants used 'memory trouble' to describe their cognitive difficulties but with further questioning often described difficulties with multi-tasking, completing tasks with multiple steps, playing board games with family, following directions/recipes, learning new computer software, mental calculations, organizing personal calendars, or planning and executing projects at home or

![Figure 7 FDG–PET in executive predominant presentations versus amnestic, visual and language predominant Alzheimer's disease phenotypes. The FDG–PET scans from participants with dysexecutive predominate presentations (n = 15) were compared to (A) amnestic predominate (n = 110), (B) visual predominate (n = 18), (C) and language predominate (n = 7) presentations of Alzheimer's disease. In each panel, the blue-black end of the spectrum encodes the t-score for a greater degree of hypometabolism in the dysexecutive phenotype with red-orange encoding the greater degree of either (A) amnestic, (B) visual or (C) language phenotype. The t-value corresponding to voxel-level P-value of FDR corrected 0.05 (A and B) or uncorrected 0.001 (C) is indicated with arrows in the colour-bar.](image)

![Figure 8 Violin scatter plots comparing variables between executive (n = 15), amnestic (n = 110), visual (n = 18) and language predominate (n = 7) Alzheimer's disease phenotypes. There was a significant main effect of phenotype for (A) age at evaluation, (B) age at disease onset and (C) hippocampal volume. There was no significant main effect of phenotype for (D) disease duration, (E) short test of mental status (STMS), (F) or cortical thickness in Alzheimer's disease signature regions. Pairwise P-values are indicated as follows: ns = 1, *p<0.05, **p<0.01, ***p<0.001, ****p<0.14.](image)
work, suggesting more executive dysfunction with prominent deficits in working memory function.

The parieto-frontal regions form an executive network (Selemon and Goldman-Rakic, 1988) that helps coordinate attentional selectivity and heavily influences working memory capacity (Rottschy et al., 2012). These networks also have a high degree of between-subject variability (Marek and Dosenbach, 2018). Impairment in executive control tests, like the Stroop Color-Word test, can be predicted by individual variation in working memory capacity and individuals with low working memory capacity had difficulty with the Stroop interference trial by two separate but complementary mechanisms (Kane and Engle, 2003). Attentional processes needed for competition resolution can only be engaged when working memory has sufficiently maintained the task/goal (i.e. ‘ignore the word and respond to color’).

The working memory network can be separated into specialized verbal working memory (phonologic loop) and visual-spatial working memory (visuospatial sketchpad) that further interact with a central executive system, which controls and regulates cognitive processes to promote the primary executive functions described above (Baddeley, 1992; Baddeley, 2012). There is conflicting evidence on lateralized contributions to these subsystems, but most evidence suggests there is bilateral activation of parieto-frontal networks in working memory tasks with specialized contributions from the left and right hemisphere depending on the task involved (Gur et al., 1994; Cabeza and Nyberg, 2000; Walter et al., 2003; Ray et al., 2008). In a study evaluating visual working memory, the bilateral intraparietal sulci were central hubs and the strength of network synchrony in this region predicted visual working memory load capacity (Palva et al., 2010). It is reasonable to hypothesize that severe dysfunction of central hubs in large-scale networks may explain the diffuse slowing seen on 14/17 EEGs in our cohort. Future studies combining electrophysiological monitoring with functional neuroimaging would help test this hypothesis.

Features of logopenic aphasia were common in our cohort (24/31) and the phonological loop (Baddeley, 1996), responsible for the rehearsal of auditory input, overlaps with the left working memory network (Jones et al., 2012). Impaired functional connectivity in the left working memory network has been associated with impaired repetition in patients with logopenic aphasia (Whitwell et al., 2015). Tau deposition also occurs in the left working memory network to a greater degree in patients with younger onset Alzheimer’s disease (Jones et al., 2017). Despite having features of logopenic aphasia with FDG-PET and tau-PET findings consistent with abnormalities in the left working memory network brain regions, the participants included in our study did not meet root criteria for primary progressive aphasia because language was not the predominant deficit and was not the initial symptom described (Gorno-Tempini et al., 2011). Our patients also displayed wide-spread cognitive dysfunction in multiple domains without a particular predilection for confrontation naming (Table 1).

Tests of auditory verbal memory—including WMS-R/III Logical memory and the AVLT—are not traditionally considered executive functioning tasks. However, working memory, particularly the phonological loop, is required for holding onto that information in the short term and organizing content for learning (i.e. encoding) and later recall. Thus, deficits in working memory would feasibly produce impairments in the ability to encode the information. Indeed, a prominent difficulty of encoding on the AVLT has been described in early-onset Alzheimer’s disease (Dickerson et al., 2017) and is consistent with working memory network impairment involving the phonologic loop (Buchsbaum et al., 2011; Stopford et al., 2012). Similar to our findings, Dickerson et al. reported relative preservation of semantic memory (i.e. confrontation naming), supporting a theoretical framework that phonologic and semantic-based networks are separable to some degree (Baddeley, 2012). Importantly, although memory retention was impaired across all measures (i.e. AVLT DR, LM II, VR II), it was not the defining feature of the presentation.

Despite evident executive dysfunction, participants in our cohort had surprisingly relatively less impairment on one measure of working memory: the combined forward and backward digit span (z-score = −1.0). Forward digit span is a simple auditory attention task (i.e. not working memory) while backward digit span is the working memory portion of the task (Kaiser et al., 2012; Egeland, 2015). We would not expect as much impairment on the forward digit span task as it measures only simple attention but backwards digit span could not be separated from the combined score using our normative data (norms not available for WMS-III or R). Therefore, difficulties in this subtest may be underrepresented in the combined score. In line with this hypothesis, Letter Number Sequencing, another working memory test, was notably worse (z-score = −1.7).

Similar to previous studies (Scheltens et al., 2017), disproportionate impairment was evident on Trails B and 13/22 participants were unable to complete the task in the allotted 300 s (an additional 6 participants did not attempt Trails B due to failure to complete Trails A), creating a floor effect. This test requires not only visual processing but also mental set shifting throughout the test. It could be argued that difficulties on Trails B may involve some disruption of the dorsal visual stream (Kravitz et al., 2011). However, participants with dysexecutive Alzheimer’s disease had normal simultanagnosia testing (Thomas et al., 2012) in over 50% of documented cases, which is a sensitive test for dorsal stream dysfunction. Additionally, participants maintained a better ability to complete Trails A (22/28 completed), which does not require task set-shifting (i.e. performance on Trails A versus Trails B in Fig. 5). All 39 participants tested with
Luria motor sequence had moderate to severe difficulties executing this sequential task, including the 12 participants who tested normally on simultanagnosia. Separate from Luria testing, praxis errors with ideomotor apraxia were also documented in 28 out of 40 cases. The large-scale parieto-frontal-basal ganglia network model of apraxia (Gross and Grossman, 2008) would be consistent with dysfunction in the working memory network needed to conceptualize, plan and execute motor tasks in these participants (Królczak et al., 2016; Matt et al., 2017). The inability to perform tasks under executive control was the main limiting factor impacting activities of daily living in this cohort, and counselling to avoid multi-tasking, environmental and emotional distractions, and emphasize strategies to facilitate sequential processing to improve daily task performance appeared to be anecdotally beneficial. While the current findings offer an important clue to the difficulties these participants experience in sequential processing, further research with standardized assessments and formal criteria for simultanagnosia, praxis and Luria testing are needed to better understand the frequency and predictive values of abnormalities for these tests in dysexecutive Alzheimer’s disease and the implications they may have on management and counselling.

Overall, the multi-domain nature of the cognitive dysfunction observed in this cohort is consistent with impaired executive functioning, particularly working memory. This type of impairment would be expected to impact cognitive performance on a wide range of mentally effortful tasks that require conscious active manipulation of abstract and/or simultaneous information streams. These deficits in working memory and cognitive flexibility have long been observed in Alzheimer’s disease in cross-sectional (Baddeley, 1986) and longitudinal studies (Baddeley et al., 1991), but in the participants described here, executive dysfunction is the core defining clinical feature. This dysfunction is related to specific disruption of brain regions involved in executive functions (Baddeley, 2003) with relative sparing of brain regions supporting other functions (e.g. memory, vision and language) (Fig. 7). These features cannot be explained by differences in disease duration or severity (Fig. 8).

Broad behavioural and personality changes were not a feature of this cohort. Apathy was the most common behavioural symptom documented at presentation (29%) and is likely related to overall executive dysfunction (Boyle et al., 2003; Barnes et al., 2015; Lohner et al., 2017). Disinhibited behavioural symptoms and inhibition problems, in general, were not a defining feature of this cohort, in contrast to abnormalities in other core executive functions (e.g. working memory and cognitive flexibility). This lack of significant dis-inhibition and personality changes may prove to be a differentiating factor between dysexecutive Alzheimer’s disease, behavioural variant frontotemporal dementia and perhaps even behavioural Alzheimer’s disease. Further studies directly comparing these syndromes will be needed to assess the predictive value of behavioural changes as they relate to the primary brain networks involved.

CSF studies demonstrated that 24% of participants had p-tau below the nominal threshold (<61 pg/ml). This threshold is based on enzyme-linked immunosorbent assay testing in clinically diagnosed Alzheimer’s disease from non-Alzheimer’s disease causes of mild cognitive impairment or dementia (Humpel, 2011; Simonsen et al., 2017). Although it is possible that some of our participants with abnormal amyloid biomarkers and low CSF p-tau levels had an additional non-Alzheimer’s disease cause of their cognitive impairment, 5 of these 12 participants underwent tau PET, all of which were unequivocally positive (mean SUVR 2.05). One of these participants also underwent an autopsy and had Alzheimer’s disease neuropathological change (Fig. 6F–J). Decreasing levels of CSF p-tau prior to and after symptom onset has been recently described in Dominantly Inherited Alzheimer’s Network (DIAN) participants (McDade et al., 2018). Amyloid plaque development is theorized to sequester detectable CSF amyloid, and through a similar mechanism, neurofibrillary tangle accumulation has been proposed to result in lower CSF p-tau values at symptom onset (McDade et al., 2018). Potentially supporting this hypothesis, participants with dysexecutive Alzheimer’s disease often presented with significantly elevated tau PET uptake suggestive of significant neurofibrillary tangle burden (Smith et al., 2016). We did not find any direct correlation with tau PET SUVR and CSF p-tau level, but this is likely complicated by lag time between the diagnostic measures (median time to tau PET after CSF was 12 months, IQR 5–20 months) and limited statistical power (n = 17). Regardless of the underlying reason, clinicians should not discount the possibility of Alzheimer’s disease as an aetiology based on the absence of elevated CSF p-tau. Future studies re-examining the accuracy of these cut-off values with regards to disease stage, the age of onset, confirmation via in vivo molecular imaging and post-mortem examination would further enhance our understanding in this regard.

Limitations of the present study include its lack of standardized clinical assessments. For example, although we found that apraxia and aphasia were highly prevalent, these were not universally documented, raising the possibility that negative or subtle symptoms went undocumented. Similarly, several participants did not undergo neuropsychological testing, EEG, genetic testing or tau PET imaging, bringing up the possibility of selection bias and a non-representative sample for participants who had these tests performed. The majority of these participants were being evaluated for second or third opinions, which may represent a selection bias that could potentially skew the prevalence of misdiagnosis and the biomarker profiles described in this report. The retrospective nature of this study also makes it difficult to know how the results impacted clinical interpretations. Although, the strikingly consistent anatomic pattern of neurodegeneration and tau
pathology in parieto-frontal brain regions key for performing tasks under executive control provide post hoc validity to the clinical construct of a progressive dysexecutive syndrome as defined here. Future prospective studies will be needed for a higher level of validation. Another limitation of this study is the lack of a comparison group recruited in the same way in which to estimate the sensitivity and specificity of our proposed criteria. However, this represents an initial effort in formulating diagnostic criteria for a progressive dysexecutive syndrome and further classification for those with aetiologies attributable to Alzheimer’s disease. Future studies with comparison groups will be necessary before consensus criteria can be derived.

Progressive dysexecutive syndrome due to Alzheimer’s disease was characterized by early, predominant and progressive cognitive dysfunction in executive tasks with less prominent behavioural changes. Whereas patients can be significantly impaired on cognitive tests, MRI demonstrates only subtle changes early on and FDG–PET is important to consider in accurately diagnosing these patients. Parietal lobe involvement is more characteristic than medial temporal lobe involvement which may appear unaffected in some patients. Falsely negative CSF p-tau levels were not uncommon, and future studies involving tau PET will be important to confirm more cases of false-negative CSF values.

Supplementary material

Supplementary material is available at Brain Communications online.

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Competing interests

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