

FEATURED ARTICLE

Pattern and implications of neurological examination findings in autosomal dominant Alzheimer disease

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[Correction added on May 28, 2022, after first online publication: An extraneous letter that was accidentally introduced during file preparation was removed from the first word in the article title.]

Abstract

Introduction: As knowledge about neurological examination findings in autosomal dominant Alzheimer disease (ADAD) is incomplete, we aimed to determine the frequency and significance of neurological examination findings in ADAD.

Methods: Frequencies of neurological examination findings were compared between symptomatic mutation carriers and non mutation carriers from the Dominantly Inherited Alzheimer Network (DIAN) to define AD neurological examination findings. AD neurological examination findings were analyzed regarding frequency, association with and predictive value regarding cognitive decline, and association with brain atrophy in symptomatic mutation carriers.

Results: AD neurological examination findings included abnormal deep tendon reflexes, gait disturbance, pathological cranial nerve examination findings, tremor, abnormal finger to nose and heel to shin testing, and compromised motor strength. The frequency of AD neurological examination findings was 65.1%. Cross-sectionally, mutation carriers with AD neurological examination findings showed a more than two-fold faster cognitive decline and had greater parieto-temporal atrophy, including hippocampal atrophy. Longitudinally, AD neurological examination findings predicted a significantly greater decline over time.

Discussion: ADAD features a distinct pattern of neurological examination findings that is useful to estimate prognosis and may inform clinical care and therapeutic trial designs.

KEYWORDS

Alzheimer disease, autosomal dominant Alzheimer disease, differential diagnosis, neurological examination, neurological examination findings, predictive value, prognosis

1 | INTRODUCTION

The neurological examination has formed the basis for the evaluation of neurological patients for over a century.¹ It is highly standardized, and the attribution of pathological findings to distinct brain regions is well established.² The neurological examination guides the process of diagnostic investigations and informs treatment decisions in a non-

invasive as well as time- and cost-effective manner.³ Physical examination, in combination with medical history, determined the correct diagnosis in approximately 40% of patients without any further diagnostic procedures in outpatient cohorts.^{4,5}

Autosomal dominant Alzheimer disease (ADAD) is a rare monogenic form of AD.⁶ ADAD shows comprehensive overlap with sporadic AD. With respect to clinical manifestation, both ADAD and

sporadic AD exhibit typical amnesic and atypical non-amnesic cognitive presentations⁷⁻¹¹ and non-cognitive clinical symptoms such as motor symptoms, seizures, and myoclonus.^{8,9,12-16} Neuropsychological characteristics include memory disturbance, visuospatial deficits, executive dysfunction, and in later stages, generalized cognitive decline in both AD variants.^{7,10} ADAD and sporadic AD share biomarker changes proposed by the amyloid hypothesis.^{6,17-19} Neuropathological findings in both AD forms comprise amyloid beta plaques and tau tangles with a higher burden including higher Braak scores in ADAD. Lewy body disease was reported in about 30% to 50% in ADAD and sporadic AD. Cerebral amyloid angiopathy is common in both disorders, with a higher severity in some ADAD mutations.²⁰⁻²² Non AD co-pathologies such as TAR DNA-binding protein 43 (TDP-43) pathology, argyrophilic grain disease, hippocampal sclerosis, and infarcts are much more common in sporadic AD.²¹ ADAD and sporadic AD differ in the mean age at clinical onset, since ADAD starts on average in the mid-40s and sporadic AD in the 70s.²³ As a result, individuals with ADAD usually lack the age-related comorbidities commonly seen in sporadic AD, for example peripheral neuropathy, orthopedic problems, falls and consecutive traumatic brain injury, and the aforementioned neuropathological non AD co-pathologies including infarcts.^{21,24-26} Since neurological examination can be influenced substantially by age and age-related comorbidities,²⁷ ADAD provides an opportunity to determine an AD-specific pattern of neurological examination findings.

We hypothesized that ADAD holds a distinct pattern of neurological examination findings (NEF) that may inform cognitive prognosis and clinical decision-making. Therefore, the aims of this study were to (1) determine the frequency of NEF in ADAD, (2) reveal a potential change in frequency over the disease course, (3) test the capacity of NEF to distinguish between mutation carriers (MC) and mutation non carriers (non MC) among mildly cognitive impaired individuals at risk, (4) analyze associations between NEF in ADAD and both cognitive performance and brain atrophy as assessed by magnetic resonance imaging (MRI), and (5) investigate the possibility of predicting cognitive decline over time based on NEF.

2 | METHODS

2.1 | Participants

We analyzed data from the observational study of the Dominantly Inherited Alzheimer Network (DIAN) that aims to investigate the clinical and biomarker course in individuals at risk for or with ADAD over time. That is, the DIAN observational study includes data from asymptomatic and symptomatic MC for ADAD and mutation-negative family members of ADAD MC (non MC). For this study, all patients with early-onset AD from the DIAN observational study at the time of data freeze 14 ($n = 118$) were evaluated. As it is a prerequisite for entering the DIAN study to be member of a family with a known ADAD mutation, no individuals with early-onset AD without ADAD mutations were included. Hence, all of the patients with early-onset AD studied here carried a mutation in either *presenilin 1* (*PSEN1*), the gene encoding

RESEARCH IN CONTEXT

1. **Systematic Review:** A comprehensive literature review in PubMed regarding neurological examination findings (NEF) in Alzheimer disease (AD) including a wide range of neurological manifestations in AD on a symptom and diagnosis level was performed (PubMed terms: "Alzheimer disease"/"autosomal dominant Alzheimer disease"/"familial Alzheimer disease" and "neurological examination findings"/"neurological findings"/"neurological symptoms"/"neurological manifestations").
2. **Interpretation:** NEF in AD are frequent and indicative of a broader affection of brain areas beyond those involved in cognition. This is associated with a poorer prognosis.
3. **Future Directions:** The knowledge about the association between the presence of non-cognitive NEF and a worse cognitive course may inform future therapeutic AD trial designs.

HIGHLIGHTS

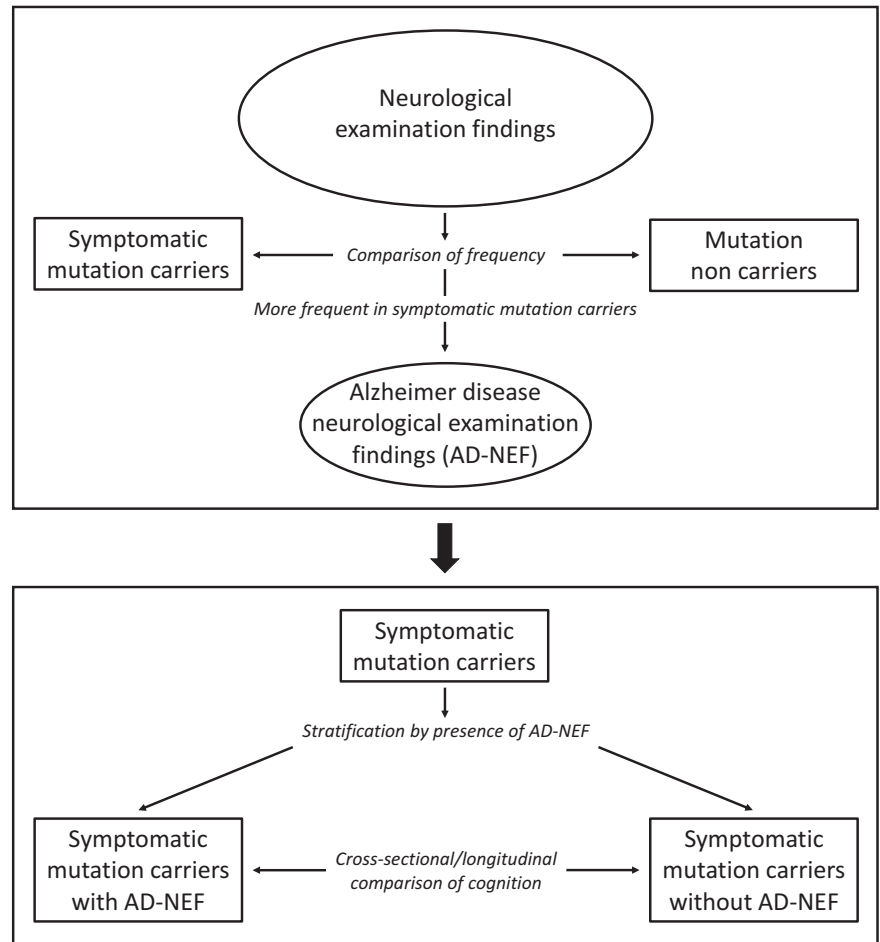
- Neurological examination findings in Alzheimer disease (AD-NEF) are frequent.
- AD-NEF are associated with a two-fold faster cognitive decline.
- Patients with AD-NEF exhibit greater parieto-temporal atrophy, including hippocampal atrophy.
- AD-NEF predict a greater cognitive decline.

for amyloid precursor protein (*APP*), or *presenilin 2* (*PSEN2*). Data were gathered at 17 study sites around the world (United States, UK, Australia, Japan, South Korea, Argentina, Spain, and Germany) between January 2008 and February 2020. Clinical data of the DIAN study participants were collected using the Uniform Data Set version 2 from the National Alzheimer's Coordinating Center (NACC-UDS2).²⁸ Clinical raters were blinded to the mutation status of the participants. The protocol of the DIAN observational study was approved by the respective institutional review boards of the study sites. The DIAN study is conducted in accordance with the declaration of Helsinki. Each study participant provided written informed consent.

2.2 | Genetic analyses

For identification of mutations in *PSEN1*, *PSEN2*, and *APP* the respective exons were amplified using polymerase chain reaction, followed by Sanger sequencing.⁶

FIGURE 1 Flow chart depicting the processes to determine Alzheimer disease neurological examinations findings (AD-NEF), of group stratifications, and of analyses performed in this study



2.3 | Neurological examination

The DIAN observational study includes a comprehensive, structured neurological examination that is completed by a trained clinical rater at each visit. The neurological examination is subdivided into 13 domains including visual impairment, auditory impairment, tremor, consciousness, cranial nerve examination findings, motor strength, finger to nose testing, heel to shin testing, sensory testing, deep tendon reflexes, plantar reflexes, gait, and other findings. Each item is rated either as absent versus present or normal versus abnormal depending on whether the respective domain label represents a pathological condition. For each rating as abnormal or present in the case of pathological conditions the study clinician may provide further details.

2.4 | Definition and classification of symptomatic ADAD

In accordance with previous studies from the DIAN,^{6,9,18} the clinical dementia rating (CDR) global score²⁹ was used to classify an individual as symptomatic (CDR global >0) or asymptomatic (CDR global = 0). For investigating the pattern of NEF across the course of ADAD, we stratified symptomatic MC by CDR global scores (groups for CDR global 0.5,

1, and 2 or 3). As numbers in the groups with CDR 2 ($n = 7$) and CDR 3 ($n = 5$) were small, these groups were taken together to form a group of MC with CDR scores of >1.

2.5 | Determination of Alzheimer disease neurological examination findings and group stratification procedures

The frequency of findings in the single subscale components of the neurological examination was compared between symptomatic MC and non MC. As both groups were relatively young (46.1 and 38.2 years, respectively) and difference in age was only 7.9 years, we did not perform statistical age matching that can cause bias itself.³⁰ For those neurological examination subscale findings that occurred more frequently in symptomatic MC than in non MC we introduced the term Alzheimer disease neurological examination findings (AD-NEF). Then, symptomatic MC were stratified by the presence of at least one AD-NEF into symptomatic MC with AD-NEF and symptomatic MC without AD-NEF. The latter stratification was done to form a cross-sectional population, that is, by the use of data from baseline visits, and a longitudinal population that included only symptomatic MC with at least the baseline visit and one follow-up visit (Figure 1).

2.6 | Calculation of disease duration

If a participant is rated as symptomatic in the DIAN observational study, the rating clinician determines the age at symptom onset by exploring the earliest progressive symptom from a predefined list of symptoms. Disease duration was calculated as the difference between the age of a participant at the time of evaluation minus her/his age at symptom onset.

2.7 | Relevant comorbidities

The data set was screened for relevant comorbidities that can influence NEF. Two participants had a history of stroke, one in the symptomatic MC group and one in the non MC group (0.8% vs 0.5%, $P = 1$). Three participants had a history of traumatic brain injury, one in the symptomatic MC group and two in the non MC group (0.8% vs 0.9%, $P = 1$). The one symptomatic mutation carrier with a history of stroke was also part of the longitudinal data set, in the symptomatic MC without AD-NEF group. The one symptomatic MC with traumatic brain injury was not part of the longitudinal data set. These participants were included in the analyses, as it was not determinable if these comorbidities actually affected NEF, were very rare, were equally distributed between groups, and in the case of stroke may be a consequence of AD-associated cerebral amyloid angiopathy.

2.8 | Magnetic resonance imaging

Structural MRI included a three-dimensional magnetization-prepared rapid acquisition with gradient echo (3D MPRAGE) sequence on 3T scanners with 1.1×1.1×1.2 mm voxel resolution. For the current analysis, we used FreeSurfer-processed (Version 6) region of interest (ROI) data (i.e., cortical thickness and subcortical volumes) in Desikan-Killiany Atlas space,³¹ provided by the DIAN neuroimaging core.

2.9 | Statistical analysis

2.9.1 | Baseline comparisons

Baseline parameters were compared using Student *t*-test for continuous variables and chi-square test or Fisher exact test for categorical variables, where appropriate.

2.9.2 | Frequencies of neurological examination findings

We used chi-square test or Fisher exact test, as appropriate, for comparison of frequencies of NEF between groups. False discovery rate correction (via Benjamini-Hochberg method) was used to account for multiple comparisons.

2.9.3 | Cross-sectional analyses

To analyze the association between the presence of AD-NEF and cognition over time, we fitted a linear mixed effects model including random intercepts with the main effects disease duration and presence/absence of AD-NEF and a disease duration*presence/absence of AD-NEF interaction term using CDR – Sum of Boxes (CDR-SB) as the outcome measure. The CDR-SB score ranges from 0 to 18, with higher values indicating worse cognitive performance. CDR-SB was chosen based on its advantages as an outcome parameter including a comprehensive assessment of cognitive performance and an almost linear decline in AD.³²

In symptomatic MC, exploratory cross-sectional structural MRI analyses were performed to determine whether the presence of AD-NEF was associated with increased gray matter atrophy determined via analyses of cortical thickness and subcortical volumes, using analyses of covariance (ANCOVA) controlling for disease duration and global Pittsburgh compound B–positron emission tomography standardized uptake value ratio (PiB-PET SUVR). Details with respect to the acquisition of PiB-PET in the DIAN observational study were described before.⁶

In addition, exploratory analyses were performed to determine whether the single AD-NEF, ataxia, or saccadic smooth pursuit eye movement were associated with specific patterns of gray matter atrophy determined via analyses of cortical thickness and subcortical volumes, using ANCOVA controlling for disease duration, PiB-PET SUVR, and for all other AD-NEF. As an indicator of ataxia, a pathological finding in either finger to nose or heel to shin testing was used.

2.9.4 | Longitudinal analyses

For investigation of a longitudinal association between AD-NEF and cognitive decline over time, that is, the rate of change in CDR-SB, a linear mixed effects model that included disease duration and presence/absence of AD-NEF at each visit as the main effects and a disease duration*presence/absence of AD-NEF interaction term with CDR-SB as the outcome parameter was used. To investigate the predictive capacity of AD-NEF regarding a future cognitive decline over time, a linear mixed effects model that included disease duration and presence/absence of AD-NEF at baseline as the main effects and a disease duration*presence/absence of AD-NEF at baseline interaction term with CDR-SB as the outcome parameter was used. The models included random slopes for each individual with variance components as the covariance matrix across random effects. For selection of the best fitting model the goodness-of-fit Akaike information criterion was used for the linear mixed effects models in this study. Linear mixed effects models were chosen for analyses because of several benefits including the ability to increase statistical power and to deal with unequal numbers of measurements or intervals.³³

Missing data were considered missing at random. All tests were performed two-sided. *P* values less than .05 were considered statistically

TABLE 1 Comparison of baseline characteristics between symptomatic mutation carriers and mutation non carriers

	Symptomatic mutation carriers (n = 118, 35.9%)	Mutation non carriers (n = 211, 64.1%)	P value
Age (years), mean (SD)	46.1 (10.0)	38.2 (11.4)	<.001
Sex (female), n (%)	56 (47.5)	87 (41.2)	.27
Education (years), mean (SD)	13.5 (3.4)	14.7 (2.9)	.001
At least one NEF, n (%)	69 (65.1)	47 (25.3)	<.001
Age at onset (years), mean (SD)	42.6 (8.8)	na	na
Disease duration (years), mean (SD)	3.7 (2.9)	na	na
CDR global, n (%)	0.5, 78 (66.1) : 1, 28 (23.7) : 2, 7 (5.9) : 3, 5 (4.2)	0, 196 (92.9) : 0.5, 15 (7.1)	<.001
CDR-SB, mean (SD)	3.8 (4.0)	0.07 (0.27)	<.001
MMSE, mean (SD)	22.5 (7.0)	29.0 (1.3)	<.001
Mutated gene, n (%)	<i>PSEN1</i> , 97 (82.2) : <i>APP</i> , 19 (16.1) : <i>PSEN2</i> , 2 (1.7)	na	na
APOE ε4 carrier, n (%)	34 (29.3)	59 (29.1)	.96

P values below .05 are italicized.

Abbreviations: SD, standard deviation; NEF, neurological examination finding; CDR, Clinical Dementia Rating; SB, Sum of Boxes; MMSE, Mini-Mental State Examination; APOE, *apolipoprotein E*.

significant. IBM SPSS Statistics Version 25 and R statistical software (Version 3.6.1) were used for statistical analyses.

2.10 | Role of the funding source

The funding source had no role in study design, in the collection, analysis, and interpretation of data, in the writing of the report, and in the decision to submit the article for publication.

3 | RESULTS

3.1 | AD-NEF

Baseline data regarding AD-NEF were available for 118 symptomatic MC and 211 non MC. Clinical and genetic parameters at baseline are listed and compared between groups in Table 1.

An abnormal neurological examination result, defined by the presence of at least one NEF, occurred more frequently in symptomatic MC compared to non MC (65.1% vs 25.3%, $P < .001$). Symptomatic MC exhibited more frequently abnormal findings in nine subdomains of the neurological examination. The highest frequency of abnormal findings showed the subdomain deep tendon reflexes (35.9% in symptomatic MC vs 3.3% in non MC, $P < .001$), followed by other findings (22.4% vs 2.7%, $P < .001$), gait (17.8% vs 2.8%, $P < .001$), cranial nerve examination findings (14.4% vs 4.7%, $P = .002$), tremor (12.7% vs 4.7%, $P = .009$), finger to nose testing (11.0% vs 0%, $P < .001$), heel to shin testing (7.7% vs 0.5%, $P = .001$), plantar reflexes (6.8% vs 1.4%, $P = .02$), and motor strength (5.1% vs 0.9%, $P = .027$). Abnormal findings in these nine subdomains of the neurological examination are referred to as AD-NEF in this article. All of these differences remained statistically significant after correction for multiple comparisons. There

were no subdomains in which abnormal neurological examination findings occurred more frequently in non MC than in symptomatic MC (Figure 2). No statistically significant differences in frequencies of AD-NEF were observed between MC with a global CDR score of 0 or in non MC (11.1% vs 14.6%; $P = .33$).

For the five most frequent AD-NEF pathological deep tendon reflexes, gait disturbance, abnormal cranial nerve examination findings, tremor, and other findings, specifications provided by the respective clinical raters were available. The most frequent findings within the respective AD-NEF were asymmetrical brisk deep tendon reflexes, reduced arm swing while walking, saccadic smooth pursuit eye movement, postural tremor, and increased muscle tone. Further specifications are depicted in Figure 3.

The ADAD MC in this study had 46 different mutations (49 *PSEN1*, 1 *PSEN2*, and 6 *APP* mutations). The single AD-NEF were compared regarding their respective frequency between single mutations using chi-square test and false discovery rate correction (via Benjamini-Hochberg method) to account for multiple comparisons. There was no difference in frequency of any AD-NEF between the single ADAD mutations.

3.2 | AD-NEFs in symptomatic MC stratified by disease stage

The frequencies of AD-NEF in symptomatic MC were analyzed by disease stage determined by CDR global scores (Figure 4). The frequency of all AD-NEF increased over the whole disease course. In all disease stages abnormal deep tendon reflexes were the most frequent finding among AD-NEF. Frequency was 33.3% at CDR 0.5, remained stable at CDR 1, and rose to 58.3% at CDR >1. Gait disturbance was present in 9.0% at CDR 0.5 and rose steadily to 28.6% at CDR 1 and to 50.0% at CDR >1. Frequency of abnormal cranial nerve examination findings

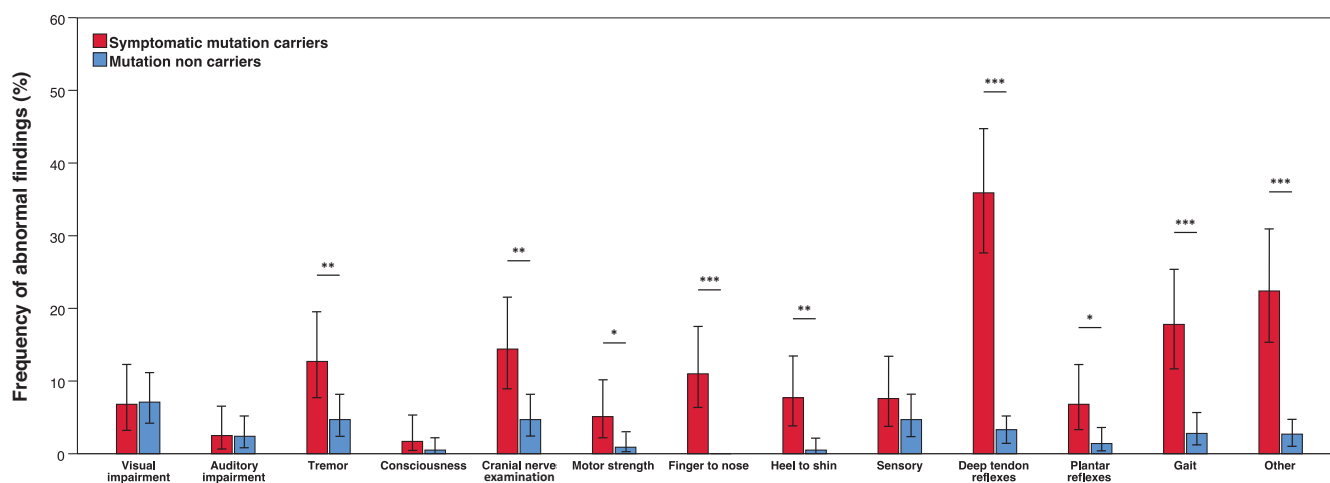


FIGURE 2 Comparisons of frequencies of neurological examinations findings between symptomatic mutation carriers and mutation non carriers. Tremor, abnormal cranial nerve examination findings, compromised motor strength, abnormal findings on finger to nose testing and heel to shin testing, pathological deep tendon reflexes, abnormal plantar reflexes, gait disturbance, and other findings were more frequent in symptomatic mutation carriers. * $P < .05$; ** $P < .01$; *** $P < .001$. Error bars represent 95% confidence intervals

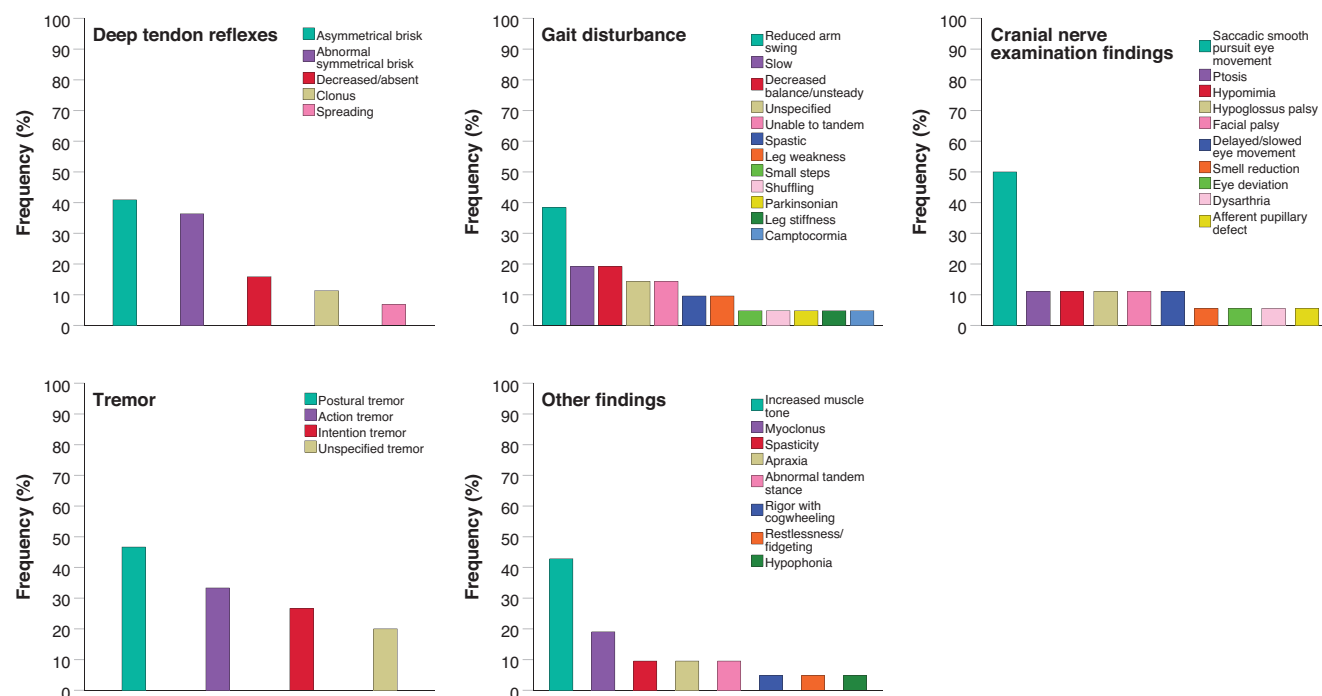


FIGURE 3 Particular signs and their frequencies within the group of the five most frequent Alzheimer disease neurological examination findings pathological deep tendon reflexes, gait disturbance, abnormal cranial nerve examination findings, tremor, and other findings. The most frequent particular signs of each of the five Alzheimer disease neurological examination findings were asymmetrical brisk deep tendon reflexes, reduced arm swing while walking, saccadic smooth pursuit eye movement, postural tremor, and increased muscle tone

was in the medium range of frequencies across disease stages: 11.5% at CDR 0.5, stayed roughly stable at CDR 1 (10.7%), and increased to 41.7% at CDR >1. Tremor occurred in 9.0% at CDR 0.5, its frequency rose slightly to 14.3% at CDR 1 and then more steeply to 33.3% at CDR >1. Abnormal finger to nose testing was found in a relatively small per-

centage of 2.6% at CDR 0.5, its frequency increased steeply to 28.6% at CDR 1, and then decreased slightly to 25.0% at CDR >1. Abnormalities in heel to shin testing exhibited the lowest frequency at CDR 0.5 (1.3%), and rose relatively steadily to 17.9% at CDR 1, and to 27.3% at CDR >1, in the medium range of frequencies of AD-NEF in the CDR 1 and

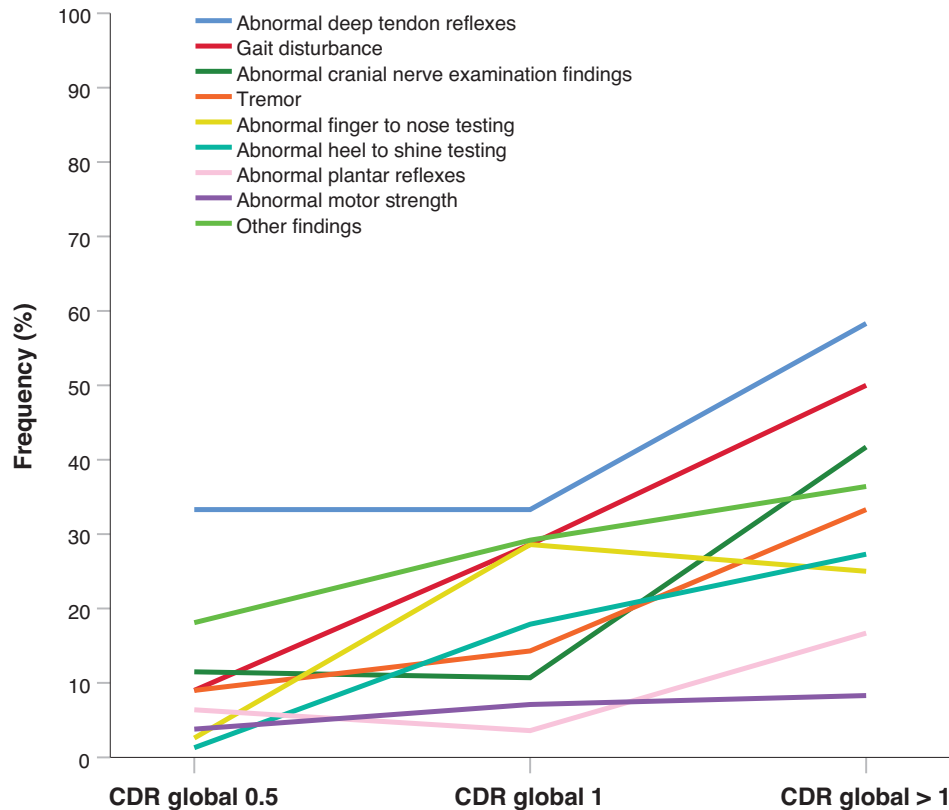


FIGURE 4 Frequencies of Alzheimer disease neurological examination findings stratified by global CDR scores. All Alzheimer disease neurological examination findings increased in frequency with CDR global stages. Abnormal deep tendon reflexes were the most frequent finding in all disease phases. Gait disturbance exhibited the steepest increase in frequency with autosomal dominant Alzheimer disease progression. Abbreviations: CDR, Clinical Dementia Rating

CDR >1 groups. Frequencies of abnormal plantar reflexes were in the lower range of frequencies through all disease stages. They were present in 6.4% at CDR 0.5, slightly declined in frequency to 3.6% at CDR 1, and rose relatively steeply to 16.7% at CDR >1. Also in the lower frequency range through all disease stages were abnormalities in motor strength. They occurred in 3.8% at CDR 0.5 and increased slightly to 7.1% at CDR 1 and to 8.3% at CDR >1.

3.3 | Association between AD-NEF and cognition

To analyze the associations between AD-NEF and cognitive performance, symptomatic MC were stratified in groups by the presence ($n = 64$) or absence ($n = 42$) of AD-NEF. Baseline clinical and genetic parameters are shown in Table 2. Symptomatic MC with AD-NEF exhibited a worse cognitive performance as assessed by CDR-SB than symptomatic MC without AD-NEF (mean CDR-SB scores: 4.32 vs 2.59, $P = .007$) while being at the same disease phase as determined by disease duration (mean disease duration: 3.9 vs 3.6 years, $P = .53$). A linear mixed effects model revealed a significant effect of the presence of AD-NEF on cognitive performance as measured by CDR-SB towards abnormal over disease duration (disease duration: estimate = 0.406, standard error = 0.186, $P = .031$; disease duration*presence of AD-

NEF interaction: estimate = 0.572, standard error = 0.221, $P = .011$). In this cross-sectional model, the decline in CDR-SB per year was 0.41 points in symptomatic MC without AD-NEF and 0.98 points in MC with AD-NEF. Symptomatic MC with AD-NEF declined significantly more, by 0.57 points on CDR-SB per year (Figure 5A).

3.4 | Association between AD-NEF and gray matter atrophy in MC

In symptomatic MC, we found that the presence of AD-NEF was associated with greater gray matter atrophy in the temporo-parietal cortex (left precuneus, left posterior cingulate, left entorhinal cortex, right superior temporal gyrus) and bilateral hippocampus at an exploratory ROI-wise alpha threshold of 0.05, controlling for disease duration and global PiB-PET SUVR (Figure 5B). At a Bonferroni-corrected alpha threshold accounting for 82 ROIs ($P < .0006$), only the left hippocampus remained significant.

The results of the exploratory analyses to determine whether the single AD-NEF, ataxia, or saccadic smooth pursuit eye movement were associated with specific patterns of gray matter atrophy are summarized in the supplementary figure. In summary, most of the single AD-NEF, ataxia, and saccadic smooth pursuit eye movement were

TABLE 2 Comparison of baseline characteristics between symptomatic mutation carriers with and without Alzheimer disease neurological examination findings

	Symptomatic MC with AD-NEF (n = 64, 60.4%)	Symptomatic MC without AD-NEF (n = 42, 39.6%)	P value
Age (years), mean (SD)	45.9 (10.3)	46.2 (8.7)	.90
Sex (female), n (%)	30 (46.9)	25 (59.5)	.20
Education (years), mean (SD)	12.9 (3.8)	14.0 (2.6)	.12
Age at onset (years), mean (SD)	42.0 (8.9)	43.1 (8.1)	.54
Disease duration (years), mean (SD)	3.98 (3.18)	3.46 (2.58)	.39
CDR global, n (%)	0.5, 40 (62.5) : 1, 15 (23.4) : 2, 5 (7.8) : 3, 4 (6.3)	0.5, 32 (76.2) : 1, 9 (21.4) : 2, 1 (2.4) : 3, 0 (0)	.20
CDR-SB, mean (SD)	4.32 (4.60)	2.62 (2.10)	.012
MMSE, mean (SD)	21.45 (7.36)	24.76 (5.22)	.008
Mutated gene, n (%)	<i>PSEN1</i> , 52 (81.3) : <i>APP</i> , 11 (17.2) : <i>PSEN2</i> ^a	<i>PSEN1</i> , 33 (78.6) : <i>APP</i> , 8 (19.0) : <i>PSEN2</i> ^a	.92
APOE ε4 carrier, n (%)	20 (31.3)	9 (21.4)	.27

P values below .05 are italicized.

Abbreviations: MC, mutation carriers; AD-NEF, Alzheimer disease neurological examination findings; SD, standard deviation; CDR, Clinical Dementia Rating; SB, Sum of Boxes; MMSE, Mini-Mental State Examination; APOE, apolipoprotein E.

^aAs there were fewer than three *PSEN2* mutation carriers in the study, no figures are shown.

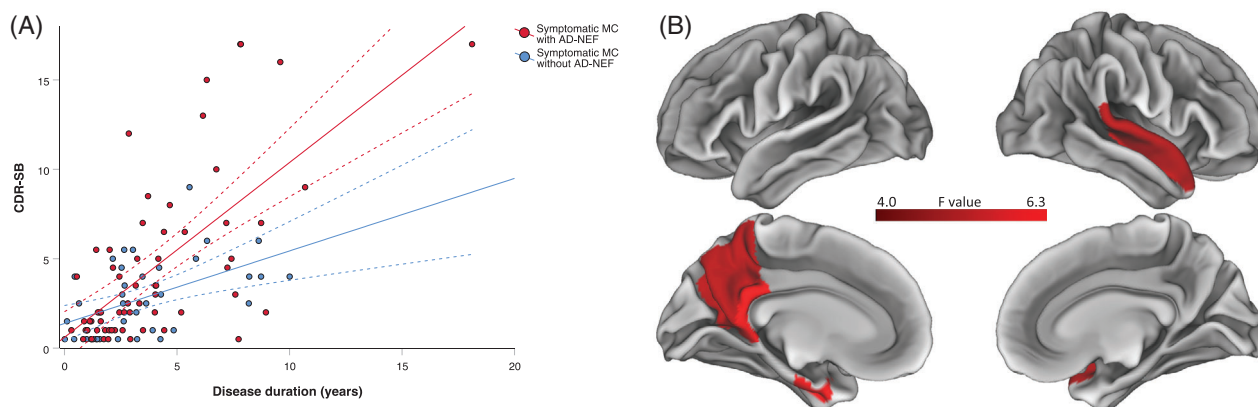


FIGURE 5 Cross-sectional associations between AD-NEF and (A) cognitive performance and (B) brain atrophy. (A) Grouped scatter plot depicting the cross-sectional relationship between CDR – Sum of Boxes scores and disease duration in symptomatic MC with and without Alzheimer disease neurological examination findings. Symptomatic MC with AD-NEF showed a significantly more pronounced decline in CDR – Sum of Boxes over the disease duration compared to symptomatic MC without AD-NEF. Dashed lines represent 95% confidence intervals. (B) Differences in brain atrophy between MC with and without AD-NEF. MC with AD-NEF showed a greater atrophy in temporo-parietal brain regions and greater bilateral hippocampal atrophy in an exploratory analysis with an alpha threshold of 0.05. After adjusting for 82 regions of interest using the Bonferroni method (resulting alpha threshold <0.0006), only the left hippocampal volume remained significantly different. Abbreviations: CDR-SB, Clinical Dementia Rating–Sum of Boxes; MC, mutation carriers; AD-NEF, Alzheimer disease neurological examination findings

associated with a fronto-temporo-parietal pattern of atrophy. There were no significant associations between any AD-NEF, ataxia, or saccadic smooth pursuit eye movement and atrophy in any subcortical region. At a Bonferroni-corrected alpha threshold accounting for 82 ROIs ($P < .0006$), no brain region remained significant.

3.5 | Longitudinal analysis and predictive value of AD-NEF regarding individual rate of cognitive decline

Longitudinal data, that is, data from the baseline visit and at least one follow-up visit of the same individual, were present for 73 symptomatic

MC with a total of 222 visits (≥ 2 visits: $n = 73$; ≥ 3 visits: $n = 39$; ≥ 4 visits: $n = 21$; ≥ 5 visits: $n = 12$; ≥ 6 visits: $n = 3$; 7 visits: $n = 1$). Mean number of visits was 3.04 (standard deviation = 1.25) and mean follow-up time 2.49 years (standard deviation = 1.63; range = 0.96–7.03 years). There was a significant difference in slopes as a function of the presence of AD-NEF at each visit and disease duration with CDR-SB as the outcome parameter (disease duration: estimate = 0.981, standard error = 0.099, $P < .001$; disease duration*presence of AD-NEF at each visit interaction: estimate = 0.343, standard error = 0.136, $P = .012$). The rate of yearly decline estimated by the model was 0.98 points on the CDR-SB score in symptomatic MC without AD-NEF compared to 1.32 points in symptomatic MC with AD-NEF. That is, symptomatic

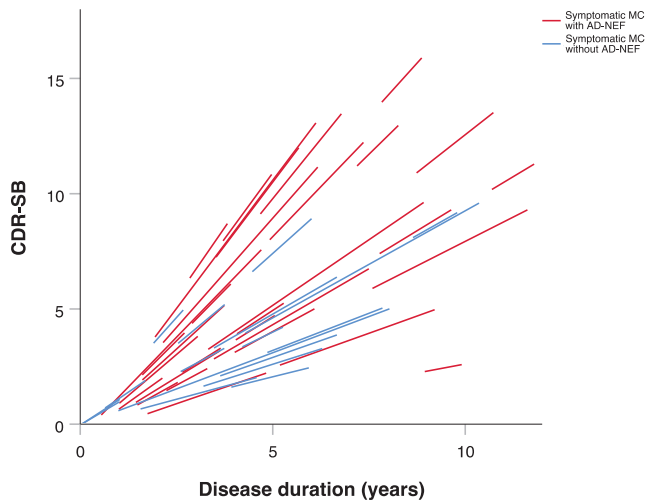


FIGURE 6 Individual linear estimates of change in Clinical Dementia Rating–Sum of Boxes (CDR-SB) score over time in symptomatic mutation carriers (MC) with and without Alzheimer disease neurological examination findings (AD-NEF). Description: Individual decline in CDR-SB score over time was significantly more pronounced in symptomatic MC with AD-NEF compared to those without. The individual linear changes in CDR-SB score were predicted by a linear mixed effects model based on longitudinal data, that is, data from symptomatic MC with at least the baseline visit and one follow-up visit. Abbreviations: MC, mutation carriers; CDR-SB, Clinical Dementia Rating–Sum of Boxes; AD-NEF, Alzheimer disease neurological examination findings

mutation carriers with AD-NEF declined significantly more, by 0.34 points per year, than symptomatic mutation carriers without AD-NEF (Figure 6). There was also a significant difference in slopes as a function of the presence of AD-NEF at baseline and disease duration with CDR-SB as the outcome parameter (disease duration: estimate = 1.020, standard error = 0.120, $P < .001$; disease duration*presence of AD-NEF at baseline interaction: estimate = 0.494, standard error = 0.211, $P = .022$). The rate of yearly decline estimated by the model was 1.02 points on the CDR-SB score in symptomatic MC without AD-NEF at baseline compared to 1.51 points in symptomatic MC with AD-NEF at baseline. That is, symptomatic mutation carriers with AD-NEF at baseline showed a significantly increased future cognitive decline, by 0.49 points on CDR-SB per year, than symptomatic mutation carriers without AD-NEF at baseline.

3.6 | Differential diagnostic significance of AD-NEF

Among individuals at risk for ADAD with a CDR global score of 0.5, that is, very mild cognitive impairment, AD-NEF were significantly more frequent in MC than in non MC (55.6% vs 26.7%; $P = .042$). The positive predictive value of AD-NEF to predict a MC status was 91%. Sensitivity was 56% and specificity was 73%.

4 | DISCUSSION

Two recently published studies, one using a European case series and the other comparing DIAN and literature data, provided insights about non-amnesic manifestations of ADAD on a symptom and diagnosis level.^{8,9} Relatively frequent symptoms were seizures, myoclonus, and behavioral or personality changes. Compared to symptoms and diagnoses, findings are less based on inductive generalization and provide the least abstract level of categorization, and therefore may provide more objective information and a high degree of cue validity.^{34,35} In the current study, a systematic investigation of single neurological findings as subscale components of a structured clinical neurological examination was performed, an approach that has not been pursued previously. Neurological examination findings in ADAD encompass pathological deep tendon reflexes, gait disturbance, cranial nerve examination findings, tremor, abnormal finger to nose and heel to shin test findings, pathological plantar reflexes, as well as compromised motor strength. Neurological examination findings in ADAD were associated with a two-fold faster cognitive decline and ADAD patients with neurological examination findings exhibited a greater parieto-temporal atrophy independent of disease duration. The presence of AD-NEF at baseline predicted an increased rate of future cognitive decline.

Knowledge about these examination findings may help clinicians to corroborate a suspected ADAD diagnosis and to distinguish from differential diagnoses of ADAD. Taking illustratively the five most frequent AD-NEF and their respective most frequent subitem (Figure 3) as the basis, a typical ADAD patient may present with asymmetrical brisk deep tendon reflexes, increased muscle tone, reduced arm swing while walking, saccadic smooth pursuit eye movements, and postural tremor.

A profile of motor symptoms measured by the Unified Parkinson Disease Rating Scale Part III was described recently in ADAD. This profile indicates that bradykinetic symptoms are the primary motor manifestation in ADAD.¹⁴ The insights about clinical neurological examination findings of this study may add further to a sharper and more comprehensive clinical picture of ADAD.

The term Alzheimer disease neurological examination findings (or AD-NEF) was introduced for those findings that were more frequent in symptomatic mutation carriers than in non mutation carriers. The frequency of AD-NEF increased with the disease stage of ADAD. This finding is in accordance with the disease phase-dependent build-up of non-cognitive symptoms in AD such as for example seizures and motor symptoms.^{9,14,15}

In at-risk individuals with mild cognitive symptoms in this study, the presence of AD-NEF was highly indicative of ADAD mutation carrier status. Since the presence of AD-NEF predicts a worse outcome in symptomatic ADAD, identifying this group early might facilitate earlier intervention and perhaps help to provide haste in confirming genetic results. The integration of knowledge of the predictive value of seizures and impaired rapid alternating hand movements regarding mutation carrier status in the cognitively presymptomatic phase of ADAD^{13,14} and of AD-NEF in cognitively symptomatic at-risk persons may help to aid patient evaluation and care throughout disease phases.

In the current study, an association between the presence of AD-NEF and poorer cognitive performance independent of the disease stage was found in ADAD patients. The exploratory MRI analysis revealed an increased temporo-parietal including hippocampal atrophy in MC with AD-NEF compared to MC without AD-NEF. A similar pattern was seen in a recent study of the spatial distribution of atrophy in ADAD patients.³³ Therefore, a potential pathophysiological explanation for the worse cognitive performance associated with AD-NEF may be a greater burden of AD-related atrophy in ADAD patients with AD-NEF independent of the disease stage.

Beyond the cross-sectional association of AD-NEF with poorer cognitive performance in ADAD patients, our intra-individual longitudinal analyses showed an association between the presence of AD-NEF and a significantly higher rate of cognitive decline over time, by approximately 35% per year on CDR-SB. Moreover, the longitudinal analysis showed that the presence of AD-NEF at baseline predicted a significantly higher rate of future decline in CDR-SB, by approximately 50% per year. The predictive capability of AD-NEF offers the opportunity to estimate prognosis and thus may add substance to patient counseling as well as to diagnostic and therapeutic strategies. Taking the stage of very mild dementia (CDR-SB 3.0-4.0) as an assumptive starting point, after 5 years patients without AD-NEF would arrive at the stage of mild dementia (CDR-SB 4.5-9.0), whereas patients with AD-NEF would be at the stage of moderate dementia (CDR-SB 9.5-15.5). After 10 years, ADAD patients without AD-NEF would exhibit moderate dementia and those with AD-NEF would suffer from severe dementia (CDR-SB 16.0-18.0).³⁶ The predictive nature of AD-NEF regarding cognitive decline over time could be explained through a potential capability of the neurological examination to detect subtle and localized AD-associated brain changes that did not yet extend to brain regions that cause cognitive decline when damaged.

Since a population with ADAD formed the basis for the analyses in this study, it is a crucial question how our findings may translate to sporadic AD. In literature, spastic paraparesis is more frequently described in ADAD than in sporadic AD. Nine of the 97 *PSEN1* mutation carriers in this study had mutations that were reported to be possibly associated with spastic paraparesis (Val261Phe, Pro264Leu, Leu271Val).³⁷ Only in one of these nine patients a bilateral spastic increase in lower limb tone was reported. Importantly, the higher age and frequency of age-related comorbidities in patients with sporadic AD that can cause abnormal NEF could challenge translatability.²⁷ Exploring for those comorbidities by thorough medical history taking including third-party anamnesis and analyses of medical files may account for these challenges and warrant translatability of the study findings to sporadic AD. However, this requires further study.

In summary, the results of our study may leverage differential diagnostic considerations by revealing neurological examination findings in symptomatic ADAD including their stage-dependent frequencies. The presence of these findings indicates mutation status in mildly cognitive impaired at-risk persons with accuracy. The association of neurological findings typical for ADAD with poor cognitive performance and their predictive value regarding increased cognitive deterioration over time may render the neurological examination suitable to contribute to

estimation of prognosis, to improve patient consultation, and to inform treatment decisions and future therapeutic trial designs.

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CONFLICTS OF INTEREST

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AUTHOR CONTRIBUTIONS

Jonathan Vöglein designed the study, wrote the manuscript, acquired, analyzed and interpreted the data, and generated the figures. Johannes Levin guided study design and concept, acquired, analyzed, and interpreted the data. Nicolai Franzmeier analyzed and visualized MRI data. Adrian Danek, John C. Morris, Marianne Dieterich, Eric McDade, Mikael Simons, Oliver Preische, Anna Hofmann, Jason Hassenstab, Tammie L. Benzinger, Anne Fagan, James M. Noble, Sarah B. Berman, Neill R. Graff-Radford, Bernardino Ghetti, Martin R. Farlow, Jasmeer P. Chhatwal, Stephen Salloway, Chengjie Xiong, Celeste M. Karch, Nigel Cairns, Richard J. Perrin, Gregory Day, Ralph Martins, Raquel Sanchez-Valle, Hiroshi Mori, Hiroyuki Shimada, Takeshi Ikeuchi, Kazushi Suzuki, Peter R. Schofield, Colin L. Masters, Alison Goate, Virginia Buckles, Nick C. Fox, Martin R. Farlow, Patricio Chrem, Ricardo Allegri, John M. Ringman, Igor Yakushev, Christoph Laske, Mathias Jucker, Günter Höglinger, and Randall J. Bateman were involved in study implementation and data collection at the respective study sites. All authors critically reviewed and revised the manuscript and figures for intellectual content.

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SUPPORTING INFORMATION

Additional supporting information may be found in the online version of the article at the publisher's website.

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