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Models of Neuroimaging, Biomarkers, and Cognitive Change in Alzheimer’s Disease
Implications for Clinical Trial Design

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Models of Neuroimaging, Biomarkers, and Cognitive Change in Alzheimer’s Disease

Implications for Clinical Trial Design

by Philip S. Insel

DOCTORAL DISSERTATION

by due permission of the Faculty of Medicine, Lund University, Sweden.
To be defended at Segerfalksalen, on 21 May 2021 at 13:00.

Faculty opponent
Michael Egan
Abstract

Objectives: Identify a window for early treatment by estimating the time course of early pathophysiological changes in Alzheimer’s disease, clarify the relationship between emerging pathology and symptom onset as well as estimate the time to clinically meaningful decline in order to inform clinical trial design.

Methods: The participants included in the analyses of the five papers were drawn from four cohorts: ADNI, AIBL, BioFINDER, A4. Repeated measures of longitudinal MRI, PET, CSF and cognitive responses were modeled using (1) mixed-effects regression with a random intercept and slope or (2) generalized least squares. Nonlinearity in longitudinal responses was captured using restricted cubic splines. Clinical trial scenarios were simulated to estimate the power to detect assumed drug effects.

Results: Clinical trials in preclinical AD are generally underpowered to detect a plausible treatment effect. Optimal composites to capture decline in the observed preclinical AD population were equal weight composites across all available cognitive and functional measures. Estimates of several major milestone events of AD progression include changes in CSF Aβ42 29 years before Aβ-positivity, an increase in regional Aβ PET deposition 15 years before, increases in tau pathology 7–8 years before, and signs of cognitive dysfunction 4–6 years before Aβ-positivity. Cognitively unimpaired Aβ+ participants approach early MCI cognitive performance levels on general cognition six years after baseline. To achieve 80% power to detect a 25% treatment effect, 2,000 participants/group for a 4-year trial and 600 participants/group for a 6-year trial are required.

Discussion: Including a large number of components in a cognitive/functional composite endpoint may smooth over aberrations in scores in a particular assessment from visit to visit within a subject, thus lowering the within-subject variance and improving signal to noise. In later stage preclinical AD, suitable power for a phase III trial can be achieved with considerably lower sample sizes while capturing both cognitive and functional change to demonstrate a clinically meaningful drug effect—both while initiating treatment in subjects who are still cognitively unimpaired. Small but meaningful increases in levels of CSF tau and temporoparietal tau are observed years before the current threshold for Aβ-positivity. In the context of secondary prevention trials, tau levels in these participants would already have been increasing for several years, likely more. These data support the use of primary prevention trials against Aβ where treatment is initiated years before the current threshold for Aβ-positivity. The separation between cognitively unimpaired participants and early MCI was just over one SD on the PACC, suggesting that one point of additional decline in Aβ+ participants compared to Aβ− participants could be taken as an approximate benchmark for clinically meaningful decline. Based on the PACC estimates, a treatment effect of 40%–50% would be required to delay the cognitive decline of a group of Aβ+ participants from reaching the one SD milestone by three years.

Key words
Models of Neuroimaging, Biomarkers, and Cognitive Change in Alzheimer’s Disease

Implications for Clinical Trial Design

by Philip S. Insel
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M’darlings Pap, Sallytoe, Claw —

95, 1062, 2700, 789…time of my life.
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Abbreviations

A4  Anti-Amyloid Treatment in Asymptomatic Alzheimer’s (study)
ADAS13 Alzheimer’s Disease Assessment Scale—Cognitive Subscale, 13 item
AIBL Australian Imaging, Biomarkers & Lifestyle (study)
AD Alzheimer’s Disease
ADNI Alzheimer’s Disease Neuroimaging Initiative
AIC Akaike Information Criterion
APOE Apolipoprotein E
Aβ Amyloid β
AVLT Auditory Verbal Learning Test
BioFINDER Biomarkers for Identifying Neurodegenerative Disorders Early and Reliably (study)
cCN Cognitively Normal Converter
CDR-SB Clinical Dementia Rating—Sum of Boxes
CI Confidence interval
CSF Cerebrospinal fluid
CU Cognitively Unimpaired
ERC Entorhinal Cortex
FAQ Functional Activities Questionnaire
FCSRT Free and Cued Selective Reminding Test
LASSO Least Absolute Shrinkage and Selection Operator
LPL Lateral Parietal Lobe
LTL Lateral Temporal Lobe
MMRM Mixed-Model Repeated Measures
MMSE Mini-Mental State Examination
MPL Medial Parietal Lobe
MRI Magnetic Resonance Imaging
<table>
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<tr>
<td>MTL</td>
<td>Medial Temporal Lobe</td>
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<tr>
<td>ng/L</td>
<td>Nanograms per Liter</td>
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<td>OR</td>
<td>Odds Ratio</td>
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<td>PACC</td>
<td>Preclinical Alzheimer’s Cognitive Composite</td>
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<td>pcCN</td>
<td>Predicted Cognitively Normal Converter</td>
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<td>PET</td>
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<td>P-tau</td>
<td>Phosphorylated Tau</td>
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<td>ROI</td>
<td>Region of Interest</td>
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<tr>
<td>sCN</td>
<td>Stable Cognitively Normal</td>
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<td>Standard Deviation</td>
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<td>SUVR</td>
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<td>T-tau</td>
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List of publications

I. Cognitive and functional changes associated with Aβ pathology and the progression to mild cognitive impairment

II. Time between milestone events in the Alzheimer's disease amyloid cascade

III. Determining clinically meaningful decline in preclinical Alzheimer disease

IV. Predicting diagnosis and cognition with 18F-AV-1451 tau PET and structural MRI in Alzheimer's disease

V. Association between apolipoprotein E ε2 vs ε4, age, and β-amyloid in adults without cognitive impairment
   Insel, Philip S., Oskar Hansson, and Niklas Mattsson-Carlgren. *JAMA Neurology* (2020).
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The separation between cognitively unimpaired participants and early MCI was just over one SD on the PACC, suggesting that one point of additional decline in Aβ+ participants compared to Aβ− participants could be taken as an approximate benchmark for clinically meaningful decline. Based on the PACC estimates, a treatment effect of 40%–50% would be required to delay the cognitive decline of a group of Aβ+ participants from reaching the one SD milestone by three years.
Background

The focus of this thesis is the improved design of clinical trials in early Alzheimer’s Disease (AD). There is no disease-modifying treatment for AD, although at the time of this writing (March 2021), for the first time, there is one phase III anti-Aβ therapy (Sevigny et al., 2016), under review for approval by the FDA.

A series of failed attempts to treat AD dementia patients with Aβ-modifying therapies led some in the field to question the amyloid cascade hypothesis and the Aβ pathway as a target for treatment. Despite significant treatment effects on biomarkers including Aβ and tau, treatments in phase III trials have failed to show significant effects on primary clinical outcomes (Doody et al., 2014; Salloway et al., 2014). These failures led to a shift toward earlier treatment in hopes of an improved likelihood of success. Attempts at earlier treatment, especially in those without cognitive impairment, poses complications in terms of trial design. Participants without symptoms of cognitive dysfunction are unlikely to decline reliably in the way that AD dementia patients would, raising questions about the trial length, sample size and outcome measurements required to detect a treatment effect. Without clinical symptoms, participants would need to be recruited based on biomarker inclusion criteria, leading to questions about the time course of the disease in terms of the relationship between accumulating Aβ and tau pathology and emerging cognitive decline.

This thesis concentrates primarily on identifying a window for early treatment by estimating the time course of early pathophysiological changes in AD, its relationship to symptom onset, and the implications for trial design features including outcome measure sensitivity, trial length and clinically meaningful decline, as well as the power to detect meaningful treatment effects.
Introduction

The papers in this thesis attempt to answer questions regarding the optimization of clinical trial design in Alzheimer’s disease — what populations should be recruited to test potential treatments, what measures should be used to evaluate change over time and test treatment effects, what is the window for treatment that may optimize the likelihood of successful treatment and what is the expected time frame for cognitive decline in relation to biomarker and imaging changes?

Paper #1 dealt primarily with which population should be recruited, and which instruments should be used to measure clinical decline over time. Directly linked to who should be recruited is when should therapeutic interventions occur — the primary aim of paper #2 was to estimate the time course of disease, especially the initial changes, and identify an optimal window for early treatment while maintaining trial feasibility and efficiency. Paper #3 also focused on the time frame of decline and characterizing the time required to reach a clinically meaningful threshold for early cognitive dysfunction. Paper #4 evaluated the relationship between several imaging measures closely associated with clinical outcomes — imaging measures that may be essential secondary or concurrent outcomes in AD clinical trials. Finally, in a slight departure, paper #5 demonstrated the potential for a drug that might mimic the biochemical properties of the APOE2 allele — a potential early treatment option for populations at high risk for AD.

The overarching theme of these papers is the estimation of the changes that occur throughout the lifespan of AD in order to identify individuals at high risk for future cognitive decline based on the earliest signals in biomarker and imaging measures. Early identification and sensitive measures of biomarker, imaging and clinical changes are essential to facilitate and optimize clinical trials in AD.

Paper #1:

Cognitive and functional changes associated with Aβ pathology and the progression to mild cognitive impairment
In 2015, at the time this paper was being developed, the Food and Drug Administration (FDA) had recently offered draft guidance to update their recommendations on primary endpoint selection in clinical trials for early-stage AD (Food and Drug Administration, 2013). With the focus of recent clinical trials on treatment in these earlier stages of AD, including prodromal AD and preclinical AD, the FDA recognized the difficulty in demonstrating drug efficacy using prior guidelines developed for trials with subjects in the dementia stage of AD (Kozauer and Katz, 2013; McKhann et al., 2011). Trial design in later stages of AD has typically included a coprimary endpoint to demonstrate efficacy on both a cognitive and a functional assessment. However, the assessment tools used in these trials have not been validated in earlier stage subjects (Snyder et al., 2014), leading the FDA to consider the use of a single primary composite endpoint that captures both cognitive and functional decline, in trials of prodromal AD subjects. Preclinical AD subjects are, by definition, cognitively unimpaired and should not have any functional impairment due to cognitive dysfunction. We hypothesize that as the target population progresses on the continuum of decline, assessing functional changes may take a more central role in demonstrating a drug effect to be clinically meaningful. However, the feasibility and value of assessing functional decline as part of a trial endpoint in a preclinical population are unknown.

Since the FDA guidance, several cognitive composites have been developed to capture the decline specific to subjects with preclinical AD, but no attempts have been made to develop combined cognitive and functional composites. The Alzheimer’s Prevention Initiative (API) has developed cognitive composites using Presenilin 1 E280A mutation carriers (Ayutyanont et al., 2014) and also cognitively unimpaired elders who progressed to MCI or AD (Langbaum et al., 2014). A third cognitive composite, to be used as the primary endpoint in the A4 trial (Sperling et al., 2014), was developed to capture decline in Aβ+ cognitively unimpaired elders (Donohue et al., 2014).

The analyses in paper #1 sought to characterize and compare the cognitive and functional decline in (1) cognitively unimpaired individuals who progress to mild cognitive impairment (MCI) and (2) cognitively unimpaired Aβ+ individuals. Identifying the cognitive and functional assessments and their weighted combinations that maximize the longitudinal decline specific to these groups may facilitate optimizing the clinical endpoints used in clinical trials of early AD and evaluate the potential role of functional assessments in cognitively unimpaired populations.
Paper #2:

Time between milestone events in the Alzheimer’s disease amyloid cascade

Previous neuropathological and biomarker data suggest that the overall time-course of AD is several decades (Li et al., 2017; Villemagne et al., 2013). In autosomal dominant AD, the estimated years to clinical onset has been used to estimate the time-course of various biomarkers in AD (Bateman et al., 2012). However, the time-course of the spread of Aβ and tau and the onset of clinical symptoms in sporadic AD is unknown.

With repeated measures of Aβ over time, the level and rate of change with respect to the key initiating AD pathology may offer a measure of disease progression in sporadic AD. The duration of amyloid positivity (chronicity) has been shown to be associated with increased tau pathology and faster cognitive decline and valuable in explaining heterogeneity in early disease progression (Koscik et al., 2020). With level and change information, the time from the threshold for significant Aβ pathology can be estimated within individuals, providing the temporal disease progression information important for evaluating biomarker trajectories. By incorporating this longitudinal information, disease progression with respect to Aβ pathology can be represented to reflect its continuous nature, resulting in a more powerful way to model the relationship between Aβ and downstream processes.

Paper #2 Sought to demonstrate the utility and predictive ability of the time-from-Aβ-positivity (TFAβ+) formulation and to evaluate the relationships between TFAβ+ and downstream biomarker and cognitive responses in order to estimate the time of the earliest signs of progression in sporadic AD. Using serial 18F-florbetapir (Aβ) PET measurements, rates of change of Aβ were estimated and used to calculate the time-from-threshold for each participant. These subject-specific estimates of the proximity to the threshold for Aβ-positivity (Aβ+) were then used to model the trajectories and temporal ordering of other key markers in AD including CSF Aβ42, regional Aβ PET, several measures of tau including CSF phosphorylated (P-tau) and total tau (T-tau), regional 18F-flortaucipir (AV-1451) tau PET, and cognition. Estimates of the time and ordering of these pathophysiological changes may facilitate the design of future prevention trials and identify a window for early treatment.
Paper #3:

Determining clinically meaningful decline in preclinical Alzheimer disease

Demonstrating that treatments are effective during the preclinical stage will require understanding the magnitude of early Aβ–related cognitive decline in cognitively unimpaired adults (Sperling et al., 2014). Defining meaningful decline will help determine the time frame for subtle cognitive changes to progress to incipient functional decline and to identify an optimal treatment window.

The association between Aβ status and cognition in preclinical AD varies widely across studies (Baker et al., 2017; Hedden et al., 2013; Insel et al., 2018; Mormino et al., 2017; Vemuri et al., 2015; Vos et al., 2014; Wirth et al., 2013), highlighting an inconsistent picture of early cognitive decline and uncertain implications for powering a trial in early AD. Understanding how sampling variation and study design features influence estimates of cognitive decline will improve the design of trials in preclinical AD.

Paper #3 Sought to harmonize several large studies in order to (1) determine the time required for a preclinical AD population to decline in a clinically meaningful way, (2) characterize how decline differs by cognitive domain, (3) update previous study design assumptions regarding sample size, power, and the required treatment effect, and (4) identify factors that modify Aβ-related decline.

Paper #4:

Predicting diagnosis and cognition with ¹⁸F-AV-1451 tau PET and structural MRI in Alzheimer’s disease

The accumulation of Aβ is assumed to lead to tau aggregation, brain atrophy, and cognitive decline (Mattsson et al., 2015; Tosun et al., 2017). However, Aβ has limited toxicity and does not typically colocalize with changes in brain structure or function. In contrast, tau spreads within and beyond regions that show atrophy in AD and correlates with cognitive decline (Villemagne et al., 2015). Tau is therefore suspected to be essential for the development of atrophy and cognitive decline in AD. This study sought to clarify the degree to which tau aggregation and atrophy are independent processes and in which brain regions tau and atrophy are most critical for development of various clinical symptoms.

Paper #4 attempted to identify brain regions where tau pathology and brain structure are most strongly associated with cognitive features of AD and as well as test for overlapping and complementary effects of tau and brain structure. We hypothesized
that an optimized measure of tau would be superior to brain structure to identify AD and cognitive impairment. However, because atrophy and cognition may also be affected by other processes than tau pathology, including Lewy body pathology, vascular pathology, and TDP-43 pathology, we hypothesized that brain structure would provide some complementary information about cognition.

Paper #5:

Association Between Apolipoprotein E ε2 vs ε4, Age, and β-Amyloid in Adults Without Cognitive Impairment

Increasing evidence suggests that the APOE genotype and its corresponding protein (apoE) affect the pathogenesis of Alzheimer’s disease (AD) through multiple biological pathways, including the differential regulation of Aβ aggregation and clearance (Liu et al., 2013; Suidan and Ramaswamy, 2019). Although the most common recent approach in AD drug discovery is to directly target the Aβ pathway, the high prevalence of APOE ε4 in AD and the ease of identifying ε4 carriers at any age make APOE pathways appealing therapeutic targets to slow Aβ accumulation (Suidan and Ramaswamy, 2019).

By mimicking the biochemical properties associated with the apoE2 isoform, it may be possible to increase the Aβ clearance that is reduced with apoE4. However, a central question is whether apoE2 remains protective in the presence of apoE4. This question has been difficult to answer, in large part because the simultaneous carriage of both the ε2 and ε4 alleles is rare—approximately 2% of the population has the ε24 genotype (Mahley, 1988).

Paper #5 sought to determine whether apoE2 remains protective in the presence of apoE4 and to evaluate how the principal risk factors for AD (age and APOE genotype) are associated with early Aβ accumulation, measured by fluroine 18–labeled (18F)-florbetapir positron emission tomography (PET).
Aims

Paper #1: Cognitive and functional changes associated with Aβ pathology and the progression to mild cognitive impairment

(1) Characterize cognitive and functional decline in
   a. cognitively unimpaired participants who progress to mild cognitive impairment
   b. cognitively unimpaired Aβ+ participants
(2) Identify and compare the cognitive and functional assessments and their weighted combinations that maximize the longitudinal decline in these groups
(3) Evaluate the potential role of functional assessments in studies of early AD.

Paper #2: Time between milestone events in the Alzheimer’s disease amyloid cascade

(1) Demonstrate the utility and predictive ability of the time-from-Aβ-positivity (TFAβ+) formulation
(2) Evaluate the relationships between TFAβ+ and downstream biomarker and cognitive responses to estimate the time of the earliest signs of progression in sporadic AD.
(3) Identify a window for early treatment using estimates of the time and ordering of pathophysiological changes.

Paper #3: Determining clinically meaningful decline in preclinical Alzheimer disease

(1) Determine the time required for a preclinical AD population to decline in a clinically meaningful way
(2) Characterize how decline differs by cognitive domain
(3) Update previous study design assumptions regarding sample size, power, and the required treatment effect
(4) Identify factors that modify Aβ-related decline.
Paper #4: Predicting diagnosis and cognition with $^{18}$F-AV-1451 tau PET and structural MRI in Alzheimer’s disease

(1) Identify regions where tau pathology and brain structure are most strongly associated with cognitive features of AD
(2) Test for overlapping and complementary effects of tau and brain structure

Paper #5: Association Between Apolipoprotein E ε2 vs ε4, Age, and β-Amyloid in Adults Without Cognitive Impairment

(1) Determine whether apoE2 remains protective in the presence of apoE4
(2) Evaluate how the principal risk factors for AD (age and APOE genotype) are associated with Aβ accumulation
Methods

Participants

The participants included in the analyses of the five papers were drawn from four cohorts:

1. The Alzheimer’s Disease Neuroimaging Initiative (ADNI) (Mueller et al., 2005)
2. The Swedish Biomarkers for Identifying Neurodegenerative Disorders Early and Reliably Study (BioFINDER) (Palmqvist et al., 2014)
3. The Australian Imaging, Biomarkers & Lifestyle (AIBL) Study (Ellis, 2009)
4. The Anti-Amyloid Treatment in Asymptomatic Alzheimer Disease Study (The A4 Study) (Sperling et al., 2020)

ADNI participants were recruited from over 50 sites across the United States and Canada (see www.adni-info.org). Participants were enrolled during multiple phases of ADNI: ADNI-1, ADNI-Go, and ADNI-2 across the spectrum of cognitive classifications including cognitively unimpaired, subjective memory complaint, mild cognitive impairment (MCI) and Alzheimer’s disease dementia. ADNI participants were included in papers #1-3.

BioFINDER participants were enrolled consecutively at three memory outpatient clinics in Sweden. BioFINDER participants were cognitively unimpaired, had mild cognitive impairment or Alzheimer’s disease dementia. BioFINDER participants were included in papers #3-4.

AIBL participants were assessed at three sites in Australia. All AIBL participants included were enrolled into the cognitively unimpaired group and were included in paper #4.

A4 participants were enrolled at 67 clinical trial sites in the US, Canada, Australia, and Japan. All A4 participants were assessed to be cognitively unimpaired and were included in paper #5.
Participants for all studies were required to have Aβ information, whether from CSF or PET, as well as completed a neuropsychological test battery. Participants were excluded if they had a major neurologic or psychiatric illness or substance abuse. ADNI participants were excluded if the screening MRI showed evidence of infection, infarction or other focal lesion. Informed written consent was obtained from all participants at each site.

Cerebrospinal Fluid

ADNI CSF samples were collected by lumbar puncture and shipped on dry ice to the ADNI Biomarker Core laboratory at the University of Pennsylvania Medical Center for long-term storage at 80 C. CSF Aβ42 was measured using the multiplex xMAP Luminex platform (Luminex Corp, Austin, TX, USA) with the research use only INNOBIA AlzBio3 kit (Fujirebio/Innogenetics, Ghent, Belgium) (Olsson et al., 2005; Shaw et al., 2009). CSF Aβ+ was defined as CSF Aβ42 < 192.

BioFINDER CSF samples were analyzed for CSF Aβ42 and Aβ40 using ELISA assays (ADx/EUROIMMUN AG, Lübeck, Germany). CSF Aβ+ was defined as CSF Aβ42/ Aβ40 < 0.1 (Janelidze et al., 2016).

Aβ Positron Emission Tomography

ADNI 18F-florbetapir PET image data were acquired 50–70 min postinjection, and images were averaged, spatially aligned, interpolated to a standard voxel size, and smoothed to a common resolution of 8 mm full width at half maximum. Methods to acquire and process ADNI 18F-florbetapir PET image data are described in (Landau et al., 2012). Full details of acquisition and analysis can be found at http://adni.loni.usc.edu/methods/. We used an a priori defined threshold for Aβ-positivity (SUVR = 1.1) (ADNI, 2012; Joshi et al., 2012) applied to the ratio of the average of the four target regions (temporal, cingulate, frontal, and parietal lobes) and the cerebellum. Aβ PET ROI outcomes were also considered (Landau and Jagust, 2015; Mormino et al., 2009), (1) the temporal lobe (middle and superior temporal lobe), (2) the parietal lobe (precuneus, supramarginal, inferior and superior parietal lobe), (3) the cingulate gyrus (isthmus, posterior, caudal and rostral anterior cingulate), (4) the frontal lobe (pars opercularis, pars triangularis, pars orbitalis, caudal/rostral middle frontal, medial/lateral orbitofrontal, frontal pole, and superior frontal lobe), and (5) a composite of regions thought to be early in accumulating Aβ (precuneus and
posterior cingulate) (Palmqvist et al., 2017). These ROIs comprise the regions included in the global composite, grouped into individual lobes plus an additional early ROI.

$^{18}$F-florbetapir ROIs were expressed as SUVRs with a cerebellar reference region.

Aβ PET imaging in the A4 Study was done using $^{18}$F-florbetapir data, which was acquired 50 to 70 minutes postinjection. Images were realigned and averaged and then spatially aligned to a standard space template. $^{18}$F-florbetapir, sampled in a global neocortical region for Aβ, was expressed as a standardized uptake value ratio (SUVR) with a cerebellar reference region (Johnson et al., 2018). β-Amyloid positivity was defined as participants with an $^{18}$F-florbetapir PET SUVR greater than or equal to 1.10 (Clark et al., 2012; Joshi et al., 2012).

Aβ PET was used in AIBL to define Aβ-positivity. Aβ-positivity was defined as $^{18}$F-florbetapir PET SUVR >1.1 (n = 72), $^{11}$C-PiB PET SUVR >1.5 (n = 201), or $^{18}$F-flutemetamol SUVR >0.62 (n = 75) (Villemagne et al., 2014).

### Tau Positron Emission Tomography

Methods to acquire and process tau ($^{18}$F-flortaucipir) PET image data in ADNI were described in (Maass et al., 2017). Six tau ROI outcomes, corrected for partial-volume, were considered: (1) the medial temporal lobe (MTL) (amygdala, entorhinal and parahippocampal cortex; from Braak stage I and III), (2) the lateral temporal lobe (LTL) (in inferior/middle/superior temporal lobe, banks of the superior temporal sulcus, transverse temporal lobe, temporal pole; from Braak stage IV and V), (3) the medial parietal lobe (MPL) (isthmus cingulate, precuneus; from Braak stage IV and V), (4) the lateral parietal lobe (LPL) (in inferior/superior parietal lobe, supramarginal; from Braak stage V), (5) frontal lobe (pars, orbitofrontal and middle/superior frontal lobe; from Braak stage V), and (6) the occipital lobe (cuneus, lingual, pericalcarine, and lateral occipital lobe; from Braak stage III, V, and VI). $^{18}$F-flortaucipir ROIs were expressed as SUVRs with an inferior cerebellar gray matter reference region. Scanner type and site were evaluated for their association with PET outcomes through covariate adjustment. Full details of PET acquisition and analysis can be found at [http://adni.loni.usc.edu/methods/](http://adni.loni.usc.edu/methods/).

Tau PET imaging in BioFINDER was done with procedures described in Smith et al., 2016. $^{18}$F-AV-1451 was synthesized at Skåne University Hospital, Lund (Hahn et al., 2017), and PET scans were performed on a GE Discovery 690 PET scanner (General Electric Medical Systems). Partial volume error correction was performed using the Geometric Transfer Method (Rousset et al., 1998) and combined with a region-based voxelwise approach (Thomas et al., 2011). FreeSurfer parcellation in MR
space of the anatomical scan was applied to processed, coregistered, and time averaged PET images to extract regional uptake values. 18F-AV-1451 standardized uptake value ratio images were based on mean uptake over 80-100 min postinjection normalized to uptake in a gray matter masked cerebellum reference region. The same FreeSurfer regions as for MRI were included for 18F-AV-1451. Besides hippocampus and amygdala, all non-neocortical structures were removed because of susceptibility to off-target binding (Smith et al., 2016). Hippocampus may also be susceptible to off-target binding because of its proximity to the choroid plexus (Lowe et al., 2016). However, we chose to include it because it is a recognized key region for structural brain changes and to facilitate comparisons between 18F-AV-1451 PET and MRI data.

Magnetic Resonance Imaging

In BioFINDER, T1-weighted MRI was performed on 3T MR scanners (Siemens Tim Trio 3T and Siemens Skyra; Siemens Medical Solutions, Erlangen, Germany), producing a high-resolution anatomical MP-RAGE image (TR5 1950 ms, TE53.4 ms, 1 mm isotropic voxels, and 178 slices). Cortical reconstruction and volumetric segmentation were performed with the FreeSurfer (v5.3) image analysis pipelines (MP-RAGE images underwent correction for intensity homogeneity (Sled et al., 1998), removal of nonbrain tissue, and segmentation into gray matter and white matter with intensity gradient and connectivity among voxels (Dale et al., 1999; Fischl et al., 2002; Fischl and Dale, 2000). Cortical thickness was measured as the distance from the gray matter/white matter boundary to the corresponding pial surface (Fischl and Dale, 2000). Reconstructed data sets were visually inspected for accuracy, and segmentation errors were corrected. Bilaterally averaged thickness measures of all available neocortical areas, plus volumes of hippocampus and amygdala, were included.

Cognitive and Functional Measures

Cognitive measures assessed in ADNI were the Mini Mental State Examination (MMSE), Alzheimer’s Disease Assessment Scale cognitive subscale, 13-item version (ADAS13), immediate and delayed memory recall from the Wechsler Memory Scale immediate and delayed Rey Auditory Verbal Learning Test, Trail Making Test parts A and B (Trails A & B), Boston Naming Test, and Category Fluency. The Clinical Dementia Rating Sum of Boxes (CDR-SB) was also assessed, which includes both cognitive and functional items, and finally the Functional Assessment Questionnaire
(FAQ), which is purely a functional assessment (Morris, 1993; Pfeffer et al., 1982; Reitan, 1958; Rey, 1958; Rosen et al., 1984; Wechsler, 1987).

Cognitive and functional measures in BioFINDER included the MMSE, immediate and delayed word list recall tests from the Alzheimer’s Disease Assessment Scale–cognitive subscale, Trail Making Test part A & B, and category (animal) fluency.

Measures assessed for AIBL participants included the MMSE, Logical Memory Delayed Recall, Digit Symbol Substitution Test, the Delayed Recall from the California Verbal Learning Test, and the CDR-SB.

A4 participants completed a neuropsychological test battery including the Preclinical Alzheimer’s Cognitive Composite (PACC) (Donohue et al., 2017, 2014) comprising the MMSE, Logical Memory Delayed Recall, Free and Cued Selective Reminding Test (FCSRT96), and the Digit Symbol Substitution Test. To calculate the PACC, the individual components were centered on their means and scaled to their standard deviations and summed, calculated using all participants. This sum was then centered on the mean and standard deviation of the sum, calculated using only the Aβ-negative group. We evaluated the FCSRT96 formulation of the FCSRT as well as the Free Recall portion of the FCRST because of evidence of their sensitivity to early Aβ-related cognitive changes. (Donohue et al., 2014; Mormino et al., 2017; Papp et al., 2017, 2015).

Modified versions of the PACC were calculated for ADNI, BioFINDER and AIBL. For ADNI, the modified PACC comprised the MMSE, Logical Memory Delayed Recall, Trail-Making Test B (Trails B), and the Delayed Word Recall from the Alzheimer’s Disease Assessment Scale–Cognitive Subscale. For AIBL, the PACC was constructed using the MMSE, Logical Memory Delayed Recall, Digit Symbol Substitution Test, and the Delayed Recall from the California Verbal Learning Test. For BioFINDER, the PACC consisted of the MMSE, Delayed Word Recall from the Alzheimer’s Disease Assessment Scale–Cognitive Subscale and Trails B.

**Statistical Methods**

Repeated measures of longitudinal imaging, biofluid and cognitive responses were modeled using (1) mixed-effects regression with a random intercept and slope or (2) generalized least squares. For development of composite cognitive and functional endpoints, numerical optimization via bound constrained optimization (Byrd et al., 1995) was used. Nonlinearity in longitudinal responses was captured using restricted cubic splines. Differences in group trajectories (frequently Aβ- vs Aβ+) over time were tested using likelihood ratio tests. P-values from multiple comparisons over many
cognitive domains or imaging ROIs were adjusted using a Hochberg correction (Hochberg, 1988). Cross-validation was used to protect against overfitting. Clinical trial scenarios were simulated and the power to detect assumed drug effects were estimated using mixed model repeated measures (MMRM) change and variance estimates.

Several analyses, especially in paper #2, focused on estimating the time course of imaging, biomarker and cognitive changes over the lifetime of AD. Each of these analyses were done in two steps. Step one involved estimating a time-from-threshold measure to place individuals on a pathological timeline. This measure used time from significant Aβ deposition (TFAβ+) as the anchoring event. TFAβ+ was estimated based on the longitudinal measures of global Aβ PET SUVR. In step two, TFAβ+ estimates were used to predict cross-sectional measures of regional tau and Aβ PET, CSF and cognitive outcomes. To demonstrate the value of the TFAβ+ measure, we did head-to-head comparisons of (i) TFAβ+ vs (ii) intercepts and slopes of longitudinal Aβ PET, modeled separately, to predict the outcomes.

Because TFAβ+ was not directly observed, in step one, linear mixed-effects models were fit to all available longitudinal global Aβ PET SUVR data to estimate subject-specific intercepts and slopes of Aβ pathology. Because Aβ slopes are unlikely to remain constant over long periods of time as subjects move toward and away from the Aβ threshold, natural splines (Hastie and Tibshirani, 1990) were used to estimate the nonlinear shape of the slopes with respect to baseline Aβ, using quantile regression (Koenker and D’Orey, 1987). Rather than modeling the mean Aβ slope with respect to baseline Aβ, quantile regression provides a separate curve for each quantile, allowing the relationship between slope and intercept to differ depending on the location in the distribution of Aβ slope. For each subject, TFAβ+ was estimated by integrating over each subject’s quantile curve from the subject’s intercept to the threshold for Aβ-positivity (PET SUVR = 1.1). For example, for a subject with a baseline SUVR of 1.2 and a slope in the 0.6 quantile, TFAβ+ was taken to be the time it would take to go from SUVR = 1.1 to 1.2, using the slope estimates from the quantile curve. For incremental changes on the x-axis (baseline SUVR), the time required to travel the incremental distance is equal to distance/rate. Using the trapezoid rule (Atkinson, 1989), TFAβ+ is the sum of these incremental times spanning SUVR = 1.1–1.2. Sensitivity analyses were done to determine the effect of varying the threshold for Aβ+. We repeated the estimation of TFAβ+ using an early threshold (SUVR 1.07) and a late threshold (SUVR 1.13).

Methods for paper #5 included estimating the individual and joint ability of 18F-AV-1451 and MRI to predict diagnosis and cognition and was done in two steps. Step one: 18F-AV-1451 and MRI composite scores. All cognitive responses (or diagnosis) were
regressed on $^{18}$F-AV-1451 and MRI, separately. For the analyses on cognition, $\text{A}\beta$-negative controls were removed, to evaluate cognition through the continuum of AD (preclinical, prodromal, and dementia stages). Each response was regressed on all included regions with $^{18}$F-AV-1451 retention levels or cortical thickness (or volume for hippocampus and amygdala), adjusting for demographics (age, sex, and years of education). The least absolute shrinkage and selection operator (LASSO) (Tibshirani, 1996) was used for model selection and to estimate regional weights to be used to form $^{18}$F-AV-1451 and MRI composites. The LASSO selects important predictors by shrinking the individual coefficients toward zero. The coefficients of covariates that do not provide additional predictive information are shrunk to zero, resulting in parsimonious and interpretable models. The LASSO is well suited to handle large numbers of highly correlated variables such as imaging regions of interest. Ten-fold cross-validation was used to tune the amount of shrinkage. Models were subsequently fit on all data using the cross-validated penalty parameter. Step two: Predictive value of $^{18}$F-AV-1451 and MRI composite scores. All responses were regressed on the composites developed in step one. The models were summarized with regression coefficients, standard errors, Wald test p-values, and the Akaike Information Criterion (AIC) (Akaike, 1974). The predictive ability of each imaging modality was summarized with classification accuracy for diagnosis and $R^2$ for cognitive responses. Ninety-five percent confidence intervals were estimated using jackknife estimated standard errors. Finally, all responses were regressed on both $^{18}$F-AV-1451 and MRI composites simultaneously to estimate the joint predictive ability of both modalities, as well as the adjusted regression coefficients, standard errors, and p-values. The reduction of the regression coefficients after adjustment was reported along with 95% confidence intervals. For prediction of diagnosis, we also present (for comparison) the accuracy of a priori selected individual regions (inferior temporal lobe for $^{18}$F-AV-1451 and hippocampal volume for MRI).
Results

Paper #1:

Cognitive and functional changes associated with Aβ pathology and the progression to mild cognitive impairment

Cohort Characteristics
In the ADNI data set, 68 subjects converted to MCI (cCN) during 7 years of follow-up while 70 subjects remained cognitively normal (sCN) throughout the same period. cCN subjects were older and had more APOE ε4 carriers compared to sCN. There were no significant differences in gender or education. We also identified a group of cognitively unimpaired subjects who were predicted to convert to MCI (pcCN, only including Aβ+ subjects).

One hundred thirty-seven Aβ+ subjects and 210 Aβ- subjects were included in the analysis. Aβ+ subjects were older, less educated, and had more APOE ε4 carriers. There was no difference in gender.

Of the 68 cCN participants, 56 had Aβ information: 31 (55.4%) were Aβ+ and 25 (44.6%) were Aβ-. Of the 70 sCN participants, 57 had Aβ information: 18 (31.6%) were Aβ+ and 39 (68.4%) were Aβ-.

Baseline cognitive/functional differences
When baseline cognitive/functional measures were compared in cCN versus sCN, cCN subjects performed worse on all 12 outcomes. There were fewer differences on baseline cognitive/functional measures in Aβ+ versus Aβ- participants.

Longitudinal change
cCN subjects worsened significantly faster on 10 of the 12 cognitive and functional outcomes compared to sCN subjects, with the exception of the Boston Naming Test and Trails A over 7 years of follow-up. The largest effect size was in the CDR-SB, and the largest effect sizes among measures without functional items were in the immediate AVLT and the ADAS13.
Aβ+ subjects worsened significantly faster on 6 of the 12 outcomes compared to Aβ- subjects. The largest effect size was in the ADAS13. cCN subjects were more likely than sCN subjects to be missing data during the course of the 7 years of follow-up (log OR = 0.82, standard error = 0.15, p < 0.001). However, sCN subjects were selected to have a minimum follow-up time of 7 years. Aβ-positivity was not associated with increased missingness (log OR = -0.04, standard error = 0.27, p = 0.87).

Composite weight distributions

The distributions of the composite weights were estimated from 1000 bootstrap samples. Composite weights were estimated separately for the 3 groups (cCN, pcCN, and Aβ+). The largest contributing outcomes in the composite for cCN versus sCN were the 2 delayed memory recall measures (delayed Logical Memory, delayed AVLT), CDR-SB, and the MMSE. Outcomes with smaller, although nonzero, positive median weights, included Category Fluency, immediate Logical Memory, Trails A, and the Boston Naming Test. When the functional measures were excluded, the delayed memory recall measures and MMSE remained the largest weighted outcomes and ADAS13 became more heavily weighted. Composite weights that maximized the separation of pcCN and sCN subjects were also estimated. Similar to the cCN composite, the main outcomes for the pcCN composite were delayed Logical Memory, CDR-SB, and MMSE, but in contrast, included the Boston Naming Test. When functional measures were excluded, the ADAS13 carried more weight.

The composites for Aβ+ versus Aβ- were heavily weighted by ADAS13, FAQ, and MMSE, as seen below.
When functional measures were excluded, ADAS13 and MMSE dominated the composites.

**Best subset components**

The best subset results were similar to the continuous optimization results. For the cCN versus sCN comparison, 5 components provided the optimal cross-validated composite, with the MMSE, delayed Logical Memory, delayed AVLT, CDR-SB, and Category Fluency selected in nearly all cross-validation folds. For the pcCN versus sCN comparison, 7 components were selected, including the MMSE, delayed Logical Memory, delayed AVLT, CDR-SB, Category Fluency, and immediate Logical Memory in nearly all folds and occasionally either ADAS13 or Trails A.

For the Aβ+ versus Aβ- comparison, 3 components were selected — the MMSE, ADAS13, and FAQ.
Power

We estimated the power to detect a 30% slowing of decline using the average out-of-sample estimates of change and variance for each composite and group, over a range of sample sizes. The composite with flat weights across all measures was the best performing composite, attaining 80% power with 375 completers/arm in a hypothetical 30-month trial. Eighty percent power was attained with 450 completers per arm using the optimized cognitive/functional composite in a hypothetical 30-month clinical trial. Sixty-five percent of power was obtained with 500 completers per arm over a 30-month trial, using a composite with cognition only. We also compared flat weight and optimized cognitive/functional composites in 48-month trials for Aβ+ pcCN subjects. They performed similarly.

With similar sample sizes as the comparisons mentioned previously, power estimates for Aβ+ subjects never exceeded 40% with any type of composite.

Paper #2:

Time between milestone events in the Alzheimer’s disease amyloid cascade

Cohort Characteristics

Two-hundred and twenty-seven CU (127 Aβ- and 100 Aβ+), 70 Aβ+ MCI and 38 Aβ+ AD participants were included in the analysis. The diagnostic groups varied by mean age, sex, years of education, and proportion of APOE ε4+. The CU- group was significantly younger than all other diagnostic groups ($p \leq 0.04$). The MCI group had a significantly smaller proportion of females than both the CU- group ($p = 0.02$) and the CU+ group ($p = 0.05$). The MCI group had significantly lower mean years of education compared to the CU- group ($p = 0.04$) and the CU+ group ($p = 0.02$). The AD group also had significantly lower mean years of education compared to the CU- group ($p = 0.01$) and the CU+ group ($p = 0.005$). The CU- group had a significantly smaller proportion of APOE ε4 carriers than all other diagnostic groups ($p < 0.003$).

Aβ PET and estimation of TFAβ+

TFAβ+ was estimated with a median of 3 (range: 1 to 5) Aβ PET scans per participant. The average time between first and last scan was 3.3 years (SD = 2.9) and the average time between scans was 2.2 years (SD = 0.8). The correlation between subject-specific random intercepts and slopes was 0.32 (0.06 to 0.55). Across diagnoses, TFAβ+ ranged from –29 to 46 years, where higher (positive) TFAβ+ values indicate more time spent
with a significant Aβ burden. The CU- group had a significantly lower mean TFAβ+ compared to all other diagnostic groups ($p < 0.001$). The CU+ group had a significantly lower mean TFAβ+ compared to the MCI group ($p < 0.001$) and the AD group ($p < 0.001$), and the MCI was significantly lower than the AD group ($p = 0.02$).

TFAβ+ estimates were not sensitive to alternative thresholds for Aβ+ beyond a shift reflecting an earlier or later threshold. When the earlier threshold (SUVR 1.07) was used rather than SUVR 1.10, TFAβ+ estimates were shifted a median of 3.2 years earlier but remained almost perfectly correlated with TFAβ+ using the SUVR 1.10 threshold ($p = 0.996$). Similarly, when the late threshold was used (SUVR 1.13), TFAβ+ estimates shifted a median of 3.1 years later, but also remained almost perfectly correlated with TFAβ+ using the SUVR 1.10 threshold ($p = 0.997$).

**TFAβ+ Performance**

TFAβ+ was highly correlated with observed time of Aβ+ ($\rho = 0.93$, 95% CI: 0.87 to 0.97, $p < 0.001$). When also including the seven participants with a subsequent negative scan after their initial positive scan, the correlation between TFAβ+ and the observed time of Aβ+ was 0.89, 95% CI: 0.80 to 0.94, $p < 0.001$. When comparing the performance of TFAβ+ versus using Aβ intercepts and slopes as separate predictors, TFAβ+ significantly outperformed separate intercepts and slopes most, but not all of the time. TFAβ+ significantly outperformed covariate only models for all outcomes. Using TFAβ+ resulted in significantly better prediction of MTL tau ($AIC_{TFAβ+} = 345.1$, $AIC_{IntSlope} = 360.2$, $AIC_{cov} = 484.8$), MPL tau ($AIC_{TFAβ+} = 467.9$, $AIC_{IntSlope} = 472.4$, $AIC_{cov} = 532.9$), occipital lobe tau ($AIC_{TFAβ+} = 272.1$, $AIC_{IntSlope} = 274.2$, $AIC_{cov} = 325.8$), CSF Aβ ($AIC_{TFAβ+} = 1825.1$, $AIC_{IntSlope} = 1827.4$, $AIC_{cov} = 1962.1$), CSF T-tau ($AIC_{TFAβ+} = 1854.4$, $AIC_{IntSlope} = 1863.4$, $AIC_{cov} = 1902.9$), MMSE ($AIC_{TFAβ+} = 1534.8$, $AIC_{IntSlope} = 1549.4$, $AIC_{cov} = 1600.4$), and the PACC ($AIC_{TFAβ+} = 1251.2$, $AIC_{IntSlope} = 1261.8$, $AIC_{cov} = 1347.4$). There was no difference between TFAβ+ and separate intercepts and slopes for LTL tau ($AIC_{TFAβ+} = 398.6$, $AIC_{IntSlope} = 399.5$, $AIC_{cov} = 471.5$) and CSF P-tau ($AIC_{TFAβ+} = 1711.3$, $AIC_{IntSlope} = 1709.6$ $AIC_{cov} = 1748.3$) and separate intercepts and slopes was significantly better than TFAβ+ in predicting frontal lobe tau ($AIC_{TFAβ+} = 194.2$, $AIC_{IntSlope} = 188.4$, $AIC_{cov} = 245.9$) and LPL tau ($AIC_{TFAβ+} = 444.9$, $AIC_{IntSlope} = 441.9$, $AIC_{cov} = 512.3$).
Regional Aβ PET

Five regional ROIs (precuneus + posterior cingulate, frontal lobe, cingulate gyrus, temporal and parietal lobes) were estimated to reach a small, but meaningful (0.2 SD) increase in SUVR between 12 and 15 years before Aβ-positivity, i.e. TFAβ+ = 0, as seen in the figure below.
At TFAβ+ = 0, all regions showed large, significant increases in SUVR (ΔSUVR ≥ 0.11, \( p \leq 0.01 \)) with the precuneus + posterior cingulate composite showing the largest increase (ΔSUVR = 0.16, \( p < 0.01 \)) and the temporal lobe showing the smallest (ΔSUVR = 0.11, \( p < 0.01 \)). Effect sizes for all regions were large (≥ 1) by the time of Aβ+. Analyses of regional Aβ PET outcomes were repeated using robust regression with robust standard errors. The robust curves are similar to the unweighted regression curves with some mild flattening in the TFAβ+ = 5 to 25 year range. The 0.2 SD change point estimates for the increase in SUVR ranged from 18 to 20 years before Aβ-positivity (compared to 12–15 years before Aβ-positivity in the main analyses). Similar to the unweighted analyses, all regions showed significance of Aβ at TFAβ+ = 0 (\( p < 0.01 \)).

**Cerebrospinal Fluid**

A 0.2 SD drop in CSF Aβ42 was estimated to occur 29 years before Aβ-positivity (TFAβ+ = −29). At TFAβ+ = 0, CSF Aβ42 showed a very large effect size (ΔAβ42 = −68 ng/L, \( p < 0.01 \), effect size = −1.99). At TFAβ+ = −2, or two years before Aβ-positivity, the population curve passes through a previously published CSF Aβ42 threshold for Aβ-positivity (192 ng/L) (Shaw et al., 2009).

A 0.2 SD increase in CSF T-tau and P-tau was estimated to occur 7–8 years before the time of Aβ-positivity (TFAβ+ = −7 and −8, respectively). At TFAβ+ = 0, significant increases of medium effect size of T-tau (ΔT-tau = 19 ng/L, \( p = 0.04 \), effect size = 0.46) and P-tau (ΔP-tau = 12 ng/L, \( p = 0.04 \), effect size = 0.47) were observed.

For the robust regression models, the change point estimate was 26 years before Aβ-positivity for the decrease in CSF Aβ, 13 years before Aβ-positivity for CSF P-tau, and 8 years before Aβ-positivity for CSF T-tau. A more substantial flattening of the curves can be seen in both CSF P-tau and T-tau for TFAβ+ > 0. The effect size for CSF T-tau at TFAβ+ = 0 remained almost identical (0.47, \( p = 0.03 \)) and the effect size for CSF P-tau increased moderately to 0.56 (\( p = 0.01 \)). The effect size for CSF Aβ42 increased to −2.52 at TFAβ+ = 0 and remained significant (\( p < 0.01 \)).

**Tau PET**

Six regional ROIs (MTL, LTL, MPL, LPL, frontal and occipital lobes) were evaluated, show in the figure below.
Five of the six regions were estimated to reach a 0.2 SD increase in SUVR 3–5 years before Aβ- positivity, with the occipital lobe reaching a 0.2 SD increase at the time of Aβ-positivity. At TFAβ+ = 0, four regions (MTL, LTL, MPL, LPL) showed significant increases in SUVR (ΔSUVR ≥ 0.14, p ≤ 0.03) with the MTL showing the largest effect size (0.36). The frontal and occipital lobes did not increase significantly by TFAβ+ = 0 (ΔSUVR = 0.09, p = 0.06 and ΔSUVR = 0.07, p = 0.13, respectively).

The robust curves show substantial flattening for TFAβ+ > 0. The robust 0.2 SD change point estimates for the increase in SUVR for the tau PET ROIs ranged from 6 to 9 years before Aβ-positivity. The significance of changes in tau PET at TFAβ+ = 0
were similar to the unweighted analyses with the exception of the frontal lobe, which increased in effect size and became statistically significant (0.44, $p = 0.02$).

**Cognition**

The MMSE showed a 0.2 SD drop six years before Aβ-positivity, followed by the PACC four years before Aβ-positivity. Neither measure decreased significantly by the time of Aβ-positivity ($\Delta$MMSE = −0.71, $p = 0.13$, effect size = −0.30; $\Delta$PACC = −0.50, $p = 0.10$, effect size = −0.32).

The robust curves show mild flattening for TFAβ+ > 0, compared to the unweighted analyses. The change point estimates for the decrease in cognitive scores was two years before Aβ-positivity for MMSE and four years before Aβ-positivity for the PACC. The robust estimate for the effect size of decrease in MMSE scores was reduced to −0.23 but became statistically significant ($p = 0.03$). The robust estimate for the effect size of decrease in PACC scores was similar (−0.30), and also became statistically significant ($p = 0.03$).

Summary curves and 0.2 SD change points for some of the earliest changing measures of each outcome type include CSF Aβ and P-tau, precuneus + posterior cingulate Aβ PET, MTL tau PET and the PACC.

**Paper #3:**

Determining clinically meaningful decline in preclinical Alzheimer disease

**Cohort characteristics**

A total of 443 cognitively healthy controls from ADNI, 348 from AIBL, and 329 from BioFINDER were included in the study. Aβ+ groups were older, had a higher frequency of APOE e4 positivity, and performed significantly worse on several cognitive tests at baseline, compared to Aβ− groups, in all cohorts.
Cognitive changes

A\(\beta\)+ participants declined significantly more on the PACC and all individual components of the PACC compared to A\(\beta\)− participants, in all 3 cohorts, with the exception of Trails B in BioFINDER (\(p = 0.08\)).

At year 4, the A\(\beta\)+ groups declined by −0.45 points on the PACC (ADNI), −0.48 points (BioFINDER), and −0.53 points (at 4.5 years, AIBL). At year 4, the A\(\beta\)− group improved 0.09 points on the PACC in ADNI and declined by −0.14 points in BioFINDER and −0.02 points in AIBL.

Clinical significance

To evaluate decline and to characterize what might be considered a clinically significant change, we compared the scores of the cognitively unimpaired participants to the baseline scores of the early MCI participants in ADNI. The mean PACC score in A\(\beta\)− and A\(\beta\)+ early MCI participants at baseline was −1.01 and −1.30, respectively. Six years after baseline, the estimated PACC score combined across cohorts of the preclinical AD groups was midway between the A\(\beta\)− and A\(\beta\)+ early MCI performance, seen in the figure below.
Similarly, the early MCI Aβ− and Aβ+ scores at baseline on the CDRSB were 1.22 and 1.38, respectively, whereas the preclinical AD groups averaged about 1.0 at 6 years.

On each of the MMSE, delayed list learning, and executive function, the cognitively normal Aβ+ groups averaged worse scores than both MCI groups by 6 years after baseline. The cognitively normal Aβ+ groups did not approach the MCI groups’ delayed logical memory scores by 6 years after baseline. Note that delayed logical memory was not available in BioFINDER.

**Power**

Using estimates of change and variance, we calculated the power for hypothetical 4- and 6-year clinical trials for each cohort, assuming a 30% dropout rate, and various sample sizes and drug effects. In 4-year trials, assuming a 25% drug effect, i.e. a 25% slowing of cognitive decline in the treatment group, the required sample size to reach
80% power was 2,000 per group for the estimate combining all cohorts. Assuming a larger effect size of 35%, the required sample size to reach 80% power was 1,000 per group on average. In 6-year trials, assuming a 25% drug effect, the required sample size to reach 80% power was about 600 per group for the estimate combining all cohorts. Assuming a 35% effect size, the required sample size to reach 80% power was 300 per group on average.

Paper #4:

Predicting diagnosis and cognition with $^{18}$F-AV-1451 tau PET and structural MRI in Alzheimer’s disease

Cohort Characteristics

One hundred twenty-seven participants including 56 CU controls, 32 patients with prodromal AD, and 39 patients with AD dementia were examined. All prodromal AD and AD dementia participants and 27 controls (preclinical AD) were Aβ+. For models of diagnosis, we compared all CU with the combined group of prodromal AD and AD dementia patients. The prodromal/dementia AD patients were younger on average than the CU (72.5 years vs. 74.7 years, p = 0.04) and had a higher proportion of apolipoprotein E ($APOE$) ε4 positivity (defined as the presence of one or two $APOE$ ε4 alleles; 79% vs. 43%, p < 0.01). There was no difference in education (11.9 years vs. 12.2 years, p = 0.76) and a borderline significant difference in sex (63% vs. 46% male, p = 0.07). For models of cognition, we included all preclinical AD, prodromal AD, and AD dementia participants (98 persons, with 54 males, average age 73.0 years, average education 11.9 years, 75% $APOE$ ε4 positivity).

$^{18}$F-AV-1451 tau PET

The $^{18}$F-AV-1451 signal was increased in prodromal AD and AD dementia in several regions throughout the temporal, parietal, frontal, and occipital lobes. The optimal $^{18}$F-AV-1451 classifier was 93% accurate in classifying AD (prodromal AD and AD dementia) versus CU (95% CI: 89% to 97%). The regions selected for classification were the amygdala, the parahippocampal gyrus, the entorhinal cortex (ERC), the fusiform cortex, and the inferior parietal lobule. The $a$ priori selected individual region inferior temporal cortex had 89% accuracy (95% CI: 80% to 98%).

Within Aβ+ participants with preclinical or clinical AD, $^{18}$F-AV-1451 was strongly associated with all cognitive responses (p < 0.001 for all responses). LASSO selected
different regions for each cognitive test. The ERC and middle temporal gyrus were selected for MMSE, shown in the figure below.

The parahippocampal gyrus and the ERC were selected for immediate recall. The amygdala, parahippocampal gyrus, temporal pole, ERC, and fusiform cortex were selected for delayed recall. The banks of the superior temporal sulcus, inferior temporal gyrus, lateral occipital cortex, and inferior parietal lobule were selected for Trail Making A. The inferior temporal gyrus, ERC, parahippocampal gyrus, and middle temporal gyrus were selected for category fluency.

**Magnetic resonance imaging**

Structural MRI was 83% accurate in classifying participants as AD (prodromal AD and AD dementia) versus CU (95% CI 68% to 98%). The main regions selected to classify diagnosis were the ERC, hippocampus, and fusiform gyrus. The *a priori* selected
individual region hippocampus had 76% accuracy (95% CI: 60% to 92%). Within Aβ+ participants, structural MRI was also strongly associated with all cognitive scores (P < .001 for all responses). The LASSO selected different regions for the cognitive tests. The ERC, the banks of the superior temporal sulcus, and inferior parietal lobule were selected for MMSE. The parahippocampal gyrus, ERC, and the inferior parietal lobule were selected for immediate recall. The hippocampus, ERC, amygdala, parahippocampal gyrus, banks of the superior temporal sulcus, and the inferior parietal lobule were selected for delayed recall. Several regions were selected for Trail Making A, with the inferior temporal gyrus, fusiform gyrus, and isthmus cingulate being the most influential regions. The ERC was selected for category fluency.

**Competing and complementary predictive information: MRI and 18F-AV-1451**

18F-AV-1451 showed strongest associations with delayed recall (R² = 0.48), followed by immediate recall (R² = 0.41), MMSE (R² = 0.36), category fluency (R² = 0.33), and Trail Making A (R² = 0.23). MRI had similar strength of associations with delayed recall (R² = 0.48), MMSE (R² = 0.35), immediate recall (R² = 0.34), category fluency (R² = 0.29), and Trail Making A (R² = 0.22). The estimates for 18F-AV-1451 were reduced between -2% (for diagnosis) and 43% (for MMSE) when adjusting for MRI. Reduction of MRI estimates ranged from 35% (for diagnosis) to 49% (for immediate recall) when adjusting for tau. AIC selected the models (ΔAIC when comparing two models > 2 favors the model with smallest AIC) with both 18F-AV-1451 and MRI to predict diagnosis and all cognitive responses.

**Paper #5:**

**Association Between Apolipoprotein E ε2 vs ε4, Age, and β-Amyloid in Adults Without Cognitive Impairment**

Of the 6943 participants who were part of the multicenter clinical trial screening visit, 4432 adults without cognitive impairment were included (2634 women [59.4%] and 1798 men [40.6%]; mean [SD] age, 71.3 [4.7] years). Individuals had mean (SD) of 16.6 (2.8) years of education, and 1512 had a positive Aβ level (34.1%).

**APOE and 18F-Florbetapir SUVR**

APOE genotype was significantly associated with 18F-florbetapir SUVR (χ² = 708.93; p < 0.001). Every APOE allele combination was significantly different from all other combinations with the exception of ε22 vs ε23 (1.02 vs 1.02; p = 0.91) and ε22 vs ε33 (1.02 vs 1.05; p = 0.43); note the small sample size of the ε22 group (n = 25). A sample
size of 272 for the ε22 group was required to detect the observed difference from the ε33 group with 80% power. Notably, the ε23 group had a significantly lower mean $^{18}$F-florbetapir SUVR compared with the ε33 group (1.02 vs 1.05; $p = 0.01$), and the ε24 group had a significantly lower mean $^{18}$F-florbetapir SUVR compared with the ε34 group (1.11 vs 1.18; $p < 0.001$). Adjusting for cardiovascular risk score did not affect the APOE genotype estimates and was not associated with $^{18}$F-florbetapir SUVR ($\beta = 0.0001$; $p = 0.98$).

**APOE, Age, and $^{18}$F-Florbetapir SUVR**

There was a significant interaction between APOE genotype and age to predict $^{18}$F-florbetapir SUVR ($p < 0.001$; ΔAIC = –26.4). The increase in $^{18}$F-florbetapir in the ε33 group was 0.006 SUVR per year (for every 1-year increase in age). Comparing each APOE group to the ε33 group, the ε22/ε23 group (combined because of sparse data over age in the ε22 group) increased significantly more slowly (0.002 SUVR per year; $p = 0.01$); the ε24 group increased similarly to the ε33 group (0.005 SUVR per year; $p = 0.73$); the ε34 group had approximately twice the rate of the ε33 group (0.012 SUVR per year; $p < 0.001$); and the ε44 group also had approximately twice the rate, although not significantly different from the ε33 group (0.011 SUVR per year; $p = 0.23$), shown in the figure below.

The ε24 group also increased at less than half the rate of the ε34 group (rate difference: 0.005 in the ε24 group vs 0.012 in the ε34 group; $p = 0.04$). There was no significant interaction between the APOE genotype and age to predict the odds of Aβ positivity ($\chi^2 = 3.94$; $p = 0.41$).

**Aβ, APOE, and the PACC**

The association between $^{18}$F-florbetapir SUV and decreasing PACC scores did not differ by APOE genotype (ΔAIC = 23.4; $p = 0.97$). Cardiovascular risk was associated with worse PACC scores ($\beta = –0.06$; $p < 0.001$) but did not affect the interaction
between age and $APOE$ genotype to predict PACC scores ($\Delta AIC = 22.9$; $p = 0.96$). There was also no difference when comparing $\varepsilon 4$ carriers to $\varepsilon 4$ noncarriers ($\Delta AIC = 4.4$; $p = 0.67$).

When adjusting for age, sex, and education but not $^{18}$F-florbetapir SUVR, the $APOE$ $\varepsilon 34$ group performed 0.08 points worse on the PACC compared with the $APOE$ $\varepsilon 33$ group ($\beta = -0.08$; $p = 0.01$). When adjusting for cardiovascular risk, all $APOE$ estimates remained similar, including the effect of $APOE$ $\varepsilon 34$ ($\beta = -0.086$; $p = 0.007$). When also adjusting for $^{18}$F-florbetapir SUVR, the effect of the $APOE$ $\varepsilon 34$ group was removed ($\beta = -0.012$; $p = 0.71$). For the $\varepsilon 4$ carriers vs noncarriers, the unadjusted difference was $-0.084$ ($p = 0.005$); after adjusting for $^{18}$F-florbetapir, the difference was $-0.006$ ($p = 0.85$), and after adjusting for $^{18}$F-florbetapir and cardiovascular scores, the difference was $-0.009$ ($p = 0.78$).
Discussion

The overall findings of the analyses included in this thesis are

(1) Clinical trials in preclinical AD, using conventional thresholds for significant Aβ burden are generally underpowered to detect a plausible treatment effect. Aβ-positive participants do show some functional decline over the course of 4 to 5 years, warranting the inclusion of functional measures to capture decline and increase power. Optimal composites to capture decline in the observed preclinical AD population were equal weight composites across all 12 cognitive and functional measures.

(2) Based on the amyloid cascade hypothesis, a relevant overarching time scale of the disease processes could be based on the development of Aβ pathology. Integrating Aβ PET level and rate of change information places each individual on a pathological timeline. Estimates of several major milestone events of AD progression include changes in CSF Aβ42 29 years before Aβ-positivity and an increase in regional Aβ PET deposition 15 years before Aβ-positivity. Using the biomarkers tested here, the first changes in CSF Aβ42 may define the onset of AD. Increases in tau pathology were estimated to occur 7–8 years before Aβ-positivity, as measured by CSF and 5 years before, as measured by PET. Signs of cognitive dysfunction occurred 4–6 years before Aβ-positivity. These findings provide a general time scale for initial changes in sporadic AD, which may inform clinical trials aimed at specific stages of the disease.

(3) Cognitively unimpaired Aβ+ participants approach early MCI cognitive performance levels on general cognition and global outcomes, delayed list recall, and executive function by six years after baseline. To achieve 80% power to detect a 25% treatment effect, 2,000 participants/group for a 4-year trial and 600 participants/group for a 6-year trial are required, using conventional definitions of elevated levels of Aβ burden.
(4) An optimized classifier that used regional $^{18}$F-AV-1451 had superior diagnostic accuracy for AD compared to brain MRI. $^{18}$F-AV-1451 and MRI had overall similar strengths of associations with cognition. Both $^{18}$F-AV-1451 and MRI contributed complementary information about cognitive impairment through the continuum from preclinical to prodromal and dementia stages of AD, with regional differences between the modalities.

(5) In a cognitively unimpaired population, $APOE \varepsilon 2$ was associated with a reduction in both the overall and the age dependent level of Aβ in the presence of $\varepsilon 4$. Large differences in levels of Aβ between $APOE$ groups were already apparent at age 65 years. The association between Aβ and decreasing global cognitive scores did not differ by $APOE$ genotype. The associated reduction in cognitive performance in $APOE \varepsilon 4$ carriers compared with noncarriers was completely mediated by Aβ.

Paper #1 demonstrates that equal weights over a fair number of cognitive and functional tests performs well in this population. The failure of the optimization to beat the equal weight composites suggests that using either continuous weights or best subset component selection results in overfitting the training sets and a subsequent reduction of test set power. Including a large number of components in a composite may smooth over aberrations in scores in a particular assessment from visit to visit within a subject, thus lowering the within-subject variance and improving signal to noise. Similarly, the equal weight composite provided the most power in Aβ+ participants, although power did not approach levels suitable for a phase III trial.

As the number of components included in the composite increases, the magnitude of change over time decreases, however, both the within-subject and between-subject errors are decreasing at a rate that overcomes the decrease in the magnitude of change, resulting in an increasing effect size. The increase in effect size plateaus in the 6-10 component range for both the converter and Aβ+ groups. The decrease in within-subject variance is clear in both groups.

We evaluated assessments available in the ADNI neuropsychological battery, although it is possible or likely that there are other measures more sensitive to decline in preclinical AD. We also did not consider item-level data from already formed composites, which may have affected the results due to carrying insensitive items along with more sensitive ones. We also make the assumption that a treatment will slow the progression of components selected for their fast decline. In reality, it is unknown which cognitive or functional components a treatment may affect and it is possible that an endpoint comprising slower progressing domains will yield more power.
The results suggest preclinical AD subjects with lower cognitive scores at baseline decline more reliably across both cognitive and functional measures compared to Aβ+ subjects without signs of subtle cognitive dysfunction. Later stage preclinical AD may represent a more feasible target population for clinical trials designed to slow cognitive decline. In this population, suitable power for a phase III trial can be achieved with considerably lower sample sizes while capturing both cognitive and functional change to demonstrate a clinically meaningful drug effect—both while initiating treatment in subjects who are still cognitively unimpaired. Multiple measures of delayed memory recall, orientation, processing speed, as well as multiple functional measures should be considered when forming a composite. Finally, when selecting measures, erring on the side of too many components may be preferable to too few.

In paper #2, small but meaningful increases in levels of CSF tau and temporoparietal tau are observed years before the current threshold for Aβ-positivity. In the context of secondary prevention trials where Aβ-positivity at current thresholds is required for study inclusion, tau levels in these participants would already have been increasing for several years, likely more. The spread of tau beyond the MTL to the parietal lobe and other regions may be a critical milestone in the progression of AD. Considering that a 0.2 SD increase in MPL tau can potentially be detected several years before Aβ-positivity, these data support the use of primary prevention trials against Aβ where treatment is initiated years before the current threshold for Aβ-positivity, if treatment efficacy relies on early intervention, prior to the development of tau pathology.

These analyses lack the power and precision to place the temporal and parietal tau regions in a particular order with confidence, but instead demonstrate that widespread tau is increasing years before Aβ-positivity. The ADNI CU, MCI and AD cohorts are also age matched. The AD patients, on average, have dementia by age 75, while the participants in the CU cohort who may eventually develop AD, are unlikely to do so for many years, possibly decades. By design, these cohorts with age matched groups are therefore on systematically different disease trajectories with respect to age. If earlier onset is associated with a more aggressive form of the disease, then the AD cohort may have the most aggressive form while the CU cohort, the least aggressive. If the developing Aβ pathology in the ADNI CU- cohort represents a less aggressive disease process compared with a more typical AD process, the estimates reported here could be conservative and biased toward later time estimates for downstream events. The ADNI MCI cohort may represent a more typical trajectory with respect to downstream events along the Aβ pathological timeline. Additionally, the change point estimates are influenced by both biological variation and measurement error, which varies from marker to marker. Change points in measures with high variability in the “normal” range and excess measurement error may require additional biological change to detect, despite an earlier, real increase in pathology.
Incorporating longitudinal information facilitates the estimation of the time-course of downstream events such as the spread of tau and the onset of subtle cognitive dysfunction. As the technology to measure AD pathology becomes more cost effective and noninvasive, such as plasma measures of Aβ or tau (Janelidze et al., 2020; Mielke et al., 2018; Palmqvist et al., 2019; Schindler et al., 2019), longitudinal evaluations in the context of trial-ready cohorts may greatly improve early diagnosis and expedite the execution of clinical trials in early AD.

To benchmark the magnitude of cognitive decline to a measure of clinical meaningfulness, in paper #3 we compared the scores of the cognitively unimpaired participants to those classified as early MCI—a group with incipient functional decline. The separation between these groups was just over one SD on the PACC, suggesting that one point of additional decline in Aβ+ participants compared to Aβ− participants could be taken as an approximate benchmark for clinically meaningful decline. Combining results across cohorts shows the average Aβ+ participant to have the same PACC score at six years post baseline as the average patient with early MCI had at baseline. Based on the PACC estimates, a treatment effect of 40%–50% would be required to delay the cognitive decline of a group of Aβ+ participants from reaching the one SD milestone by three years.

Delaying the cognitive decline equivalent to the level of the average early MCI patient by three years may be a clinically meaningful treatment effect. But 40%–50% is a large treatment effect and highlights the difficulties in preclinical AD trial design. In order to reliably achieve 80% power for a modest, real-world effect size of 20%–30%, investors in AD research for therapeutics development will have to prepare to support larger and longer trials than are currently envisaged.

These analyses show that large sample sizes and sufficiently long follow-up times result in consistent estimates of decline in preclinical AD. Despite substantial design and sampling differences, these results support the potential for internationally conducted clinical trials in preclinical AD. However, it is likely that designers of preclinical AD treatment trials will have to prepare for larger and longer trials than are currently considered.

The findings of paper #4 suggest that although tau is the more critical measure between 18F-AV-1451 tau PET and structural brain MRI, both measures capture partly unique information that is relevant for the clinical deterioration in AD. The selected regions were mainly temporal lobe regions, where tau pathology presumably occurs in early stages of AD (ERC, amygdala, fusiform, and the parahippocampal gyrus), but also the inferior parietal lobule, which presumably is involved in later stages of the disease (Cho et al., 2016).

The optimal MRI-based classifier achieved lower accuracy and partly included similar regions as the 18FAV-1451 classifier (ERC, fusiform) plus the hippocampus, the
banks of the superior temporal sulcus, and the inferior parietal lobule. The classification accuracy for the model including both $^{18}$F-AV-1451 and MRI did not improve over $^{18}$F-AV-1451 alone, indicating that brain structure contributes little beyond tau to the identification of patients with AD.

We conclude that $^{18}$F-AV-1451 tau PET was strongly associated with AD diagnosis, with stronger associations than structural MRI. However, both $^{18}$F-AV-1451 and structural MRI were independently associated with cognitive impairment, across the entire disease continuum from preclinical to prodromal and dementia stages of AD.

In paper #5, the separation between the $\varepsilon24$ and $\varepsilon34$ groups in terms of Aβ levels becomes clear as the groups approach 70 years of age. The reduced levels of Aβ in $\varepsilon24$ compared with $\varepsilon34$ participants shown here may be one of the primary drivers behind the protective effect of the $\varepsilon2$ allele against AD dementia, shown previously in a large case-control study (Reiman et al., 2020). The $\varepsilon24$ group demonstrated an associated reduced risk of AD dementia (odds ratio, 2.68 [95%CI, 1.65-4.36]) compared with the $\varepsilon34$ group (odds ratio, 6.13 [95% CI, 5.08-7.41]) when comparing both groups with $\varepsilon33$ participants. However, the presence of the $\varepsilon2$ allele does not completely protect against Aβ positivity, as 16% of $\varepsilon2$ homozygotes in the A4 Study had positive Aβ levels, nor does it completely protect against AD dementia, as 5 of 24 $\varepsilon2$ homozygotes had a neuropathologically confirmed AD dementia diagnosis (Reiman et al., 2020). Although the APOE genotype is one of the strongest risk factors for AD, it does not determine Aβ accumulation or cognitive decline.

The APOE $\varepsilon34$ and APOE $\varepsilon44$ groups had mean SUVRs of approximately 1.10 and 1.25 at age 65, whereas the $\varepsilon33$ and $\varepsilon23$ groups had mean SUVRs near 1.0. APOE $\varepsilon4$ carrier longitudinal rates of global $^{18}$F-florbetapir change have been estimated to be 0.0044 SUVR per year in Aβ-negative individuals and 0.0126 SUVR per year in Aβ-positive individuals (Lim and Mormino, 2017), suggesting that it would take decades for the APOE $\varepsilon4$ carrier groups to reach the Aβ levels observed at age 65 years in this study. This coincides with the estimated prevalence of Aβ positivity in $\varepsilon44$ individuals between 25% and 30% at age 45 years and 10% in $\varepsilon34$ individuals at age 50 years (Jansen et al., 2018). With Aβ positivity already observable in some individuals in their 40s, the gradual accumulation likely begins much earlier. Indeed, reductions of cerebrospinal fluid Aβ have been observed in $\varepsilon44$ carriers in their 20s (Lautner et al., 2017). A protein-modifying treatment mimicking the protective effect of the $\varepsilon2$ allele against Aβ accumulation may be most effective before significant Aβ deposition. The age at which such a treatment should be initiated would vary greatly by APOE genotype and individual, but if done safely, it could be used to slow Aβ accumulation in early middle age for those at highest risk.
This study’s findings suggest that the protective effect of carrying an ε2 allele in the presence of an ε4 allele offers potential for a treatment that attempts to mimic this protective outcome in order to facilitate Aβ clearance in ε4 carriers. Such a treatment strategy is appealing, as ε4 carriers make up 67% of patients with AD dementia, and it could represent an early treatment option, as many ε4 carriers begin to accumulate Aβ in early middle age. If the goal is to interfere early in the disease process before activation of downstream pathways, AD prevention trials may consider targeting much younger people before the accumulation of high or even intermediate levels of Aβ develop.
Conclusion

Results from phase III trials of potential disease-modifying treatments in AD continue to be almost strictly negative. However, clinical trials in AD look different today compared with the trials of ten years ago (Egan et al., 2019). Imaging and biomarker research over the last decade has led to substantial gains in the understanding of the progression of AD. This understanding has transformed clinical trial design. Previously, it was common for participants to be recruited into trials without inclusion criteria regarding Aβ pathology. For anti-Aβ treatment trials, this meant that many participants would not even harbor the pathology the treatment was targeting.

Longitudinal imaging and biomarker studies over the last decade have clarified the time course of the earliest changes in Aβ and tau, further demonstrating how late the dementia stage is in the disease process and how difficult it might be to halt or reverse cognitive decline once this stage has begun. There are currently several ongoing treatment trials targeting early biomarker changes in participants without cognitive impairment, using the latest imaging and biomarker technology to monitor the progression of early AD pathology (Mintun et al., 2021).

These advances in imaging and biomarker research have facilitated clinical trials in earlier patient populations, identified a potentially optimal treatment window, and may be the key to accelerating drug discovery in AD.
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Cognitive and functional changes associated with Aβ pathology and the progression to mild cognitive impairment

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Abstract

Cognitively-normal people with evidence of β-amyloid (Aβ) pathology and subtle cognitive dysfunction are believed to be at high risk for progression to mild cognitive impairment due to Alzheimer’s disease (AD). Clinical trials in later stages of AD typically include a coprimary endpoint to demonstrate efficacy on both cognitive and functional assessments. Recent trials focus on cognitively-normal people, but functional decline has not been explored for trial designs in this group. The goal of this study was therefore to characterize cognitive and functional decline in (1) cognitively-normal people converting to mild cognitive impairment (MCI) and (2) cognitively-normal Aβ-positive (Aβ+) people. Specifically, we sought to identify and compare the cognitive and functional assessments and their weighted combinations that maximize the longitudinal decline specific to these 2 groups. We studied 68 people who converted from normal cognition to MCI and 70 nonconverters, as well as 137 Aβ+ and 210 Aβ- amyloid-negative cognitively-normal people. We used bootstrap aggregation and cross-validated mixed-models to estimate the distribution of weights applied to cognitive and functional outcomes to form composites. We also evaluated best subset optimization. Using optimized composites, we estimated statistical power for a variety of clinical trial scenarios. Overall, 55.4% of cognitively-normal to MCI converters were Aβ+. Large gains in power estimates were obtained when requiring participants to have both subtle cognitive dysfunction and Aβ pathology compared with requiring Aβ pathology alone. Additional power resulted when including functional as well as cognitive outcomes as part of the composite. Composites formed by applying equal weights to all measures provided the highest estimates of cross-validated power, although similar to both continuous weight optimization and best subset optimization. Using a composite to detect a 30% slowing of decline, 80% power was obtained for predicted Aβ+ converters with 375 completers/arm for a 30-month trial using a combination of cognitive/functional measures. In the Aβ+ group, power to approach levels suitable for a phase III clinical trial would require considerably larger sample sizes. Composites incorporating both cognitive and functional measures may substantially increase the power of a trial in a preclinical (Aβ+) AD population with subtle evidence of cognitive dysfunction.

1. Introduction

Accumulating evidence from Alzheimer’s disease (AD) biomarker studies suggests β-amyloid (Aβ) deposition may occur decades before the diagnosis of clinical dementia (Morris, 2005). Anti-Aβ treatments are thought to have a higher likelihood of slowing progression if administered at the earliest signs of the...
pathological cascade, before substantial neurodegeneration and other downstream effects of Aβ deposition (Sperling et al., 2011b). Classification of Alzheimer’s disease into progressive stages has helped to organize the current thinking about the emergence of subtle clinical symptoms and the development of cognitive and functional impairment during the continuum of disease progression (Sperling et al., 2011a). The initial stages of preclinical AD are defined by amyloidosis and neurodegeneration. The final preclinical stage also includes some evidence of subtle cognitive dysfunction, although below levels of cognitive and functional impairment required to meet criteria for mild cognitive impairment (MCI) due to AD. As the disease progresses into MCI and dementia, cognitive and functional deficits may be observed. Identifying the biomarkers and clinical assessments that can predict and monitor the progression from the early stages of AD to more advanced disease will help to elucidate the disease process and inform clinical trial design (Insler et al., 2015). Here we sought to determine the optimal combination of cognitive and functional measures to track disease progression in cognitively-normal people progressing to MCI, and of Aβ-positive (Aβ+) cognitively-normal people. Composite endpoints comprising both cognitive and functional measures are currently being used in clinical trials of MCI populations (Ard et al., 2015; Raghavan et al., 2013; Wang et al., 2016). Here we consider the inclusion of functional measures in the endpoint for clinical trials in preclinical AD.

The Food and Drug Administration (FDA) recently offered draft guidance to update their recommendations on primary endpoint selection in clinical trials for early-stage AD (US Dept of Health and Human Services, 2015). With the focus of recent clinical trials on treatment in these earlier stages of AD, including prodromal AD and preclinical AD, the FDA recognized the difficulty in demonstrating drug efficacy using prior guidelines developed for trials with subjects in the dementia stage of AD (Kozauer et al., 2013; McKhann et al., 2011). Trial design in later stages of AD has typically included a coprimary endpoint to demonstrate efficacy on both a cognitive and a functional assessment. However, the assessment tools used in these trials have not been validated in earlier stage subjects (Snyder et al., 2014), leading the FDA to acknowledge the complexity of defining an endpoint that captures both cognitive and functional decline, in trials of prodromal AD subjects. Preclinical AD subjects are, by definition, cognitively-normal and should not have any functional impairment due to cognitive dysfunction. We hypothesize that as the target population progresses on the continuum of decline, assessing functional changes may take on a more central role in demonstrating a drug effect to be clinically meaningful. However, the feasibility and value of assessing functional decline as part of a trial endpoint in a preclinical population are unknown.

Since the FDA guidance, several cognitive composites have been developed to capture the decline specific to subjects with preclinical AD, but no attempts have been made to develop combined cognitive and functional composites. The Alzheimer’s Prevention Initiative (API) has developed cognitive composites using Presenilin 1 E280A mutation carriers (Auytanont et al., 2014) and also cognitively-normal elders who converted to MCI or AD (Langbaum et al., 2014). A third cognitive composite, to be used as the primary endpoint in the A4 trial (Sperling et al., 2014), was developed to capture decline in Aβ+ cognitively-normal elders (Donohue et al., 2014), and selected individual components based on a literature review. Functional assessments were not evaluated in the API or the A4 composites.

The aim of this study was to identify and compare the cognitive and/or functional assessments and their weighted combinations that maximize the longitudinal decline specific to (1) cognitively-normal to MCI converters (cCN); and (2) cognitively-normal Aβ+ subjects. Conversion status is not known at the beginning of the study, and thus, power estimates based on subjects’ true conversion status would not be useful to inform a clinical trial. Therefore, to reflect a realistic modern trial scenario, subjects who were both predicted to convert using information available at baseline and were also Aβ+ (pCN), were used to estimate clinical trial power. Using the battery of assessments from the Alzheimer’s Disease Neuroimaging Initiative (ADNI), we sought to characterize the importance of each cognitive and functional assessment in our 3 groups (cCN, pCN, and Aβ+) as well as provide cross-validated estimates of power when using the composites in clinical trial scenarios.

2. Material and methods

2.1. Participants

Data were obtained from the ADNI database (adni.loni.usc.edu). ADNI is the result of efforts of many co-investigators and participants have been recruited from over 50 sites across the United States and Canada (see www.adni-info.org). The population in this study included ADNI-1 and ADNI-2 participants enrolled into the cognitively-normal or subjective memory complaint cohorts, were tested for cerebrospinal fluid (CSF) biomarkers or 18F-florbetapir positron emission tomography (PET), and were followed longitudinally for neuropsychological testing.

2.2. Cerebrospinal fluid biomarker concentrations

Each CSF sample was collected by lumbar puncture and shipped on dry ice to the ADNI Biomarker Core laboratory at the University of Pennsylvania Medical Center for long-term storage at ~80 °C. CSF Aβ42 was measured using the multiplex xMAP Luminex platform (Luminex Corp, Austin, TX, USA) with the research use only INNOMIA AlzBio3 kit (Fujirebio/Innogenetics, Ghent, Belgium) (Olsson et al., 2005; Shaw et al., 2009).

2.3. Florbetapir PET

Methods to acquire and process ADNI florbetapir PET image data were described previously (Landau et al., 2012). Full details of acquisition and analysis can be found at http://adni.loni.usc.edu/methods/.

2.4. Cognitive and functional outcomes

Cognitive measures assessed were the Mini-Mental State Examination (MMSE), Alzheimer’s Disease Assessment Scale—cognitive subscale, 13-item version (ADAS13), immediate and delayed memory recall from the Wechsler Memory Scale (iMemory, dMemory), Trail Making Test parts A and B (Trails A & B), Boston Naming Test, and Category Fluency. The Clinical Dementia Rating Sum of Boxes (CDR-SB) was also assessed, which includes both cognitive and functional items, and finally the Functional Assessment Questionnaire (FAQ), which is purely a functional assessment (Kaplan et al., 1982; Morris, 1993; Pfeffer et al., 1982; Reitan, 1958; Rey, 1964; Rosen et al., 1984; Wechsler, 1987).

2.5. Statistical analysis

This study included 3 main sets of analyses. The first was a comparison of normal participants who converted to a diagnosis of MCI (cCN) during 7 years of follow-up versus stable cognitively-normal (sCN) participants, during the same period. Follow-up on
converters continued beyond diagnosis of MCI. The second analysis was a comparison of participants predicted to convert to MCI (pcCN, only including Aβ+ subjects) versus the β-amyloid-negative (Aβ−) participants from the scCN group. The third analysis was a comparison of Aβ+ versus Aβ− participants, irrespective of conversion information. There was considerable overlap among these groups. The cCN or sCN participants that also had Aβ information (n = 56 from the cCN group, and n = 57 from the scCN group) were also included in either the Aβ− or Aβ+ groups. This is described further in the results section. All pcCN participants were included in the Aβ+ group.

Aβ+ was defined as flurbirapir PET SUV > 1.10 at any point during follow-up (Landau et al., 2012). Subjects without flurbirapir PET were considered Aβ− if CSF Aβ42 < 192 ng/L (Shaw et al., 2009).

In each of the 3 groups, we compared 2 types of optimization: the first allowed continuous weights for each component while the second was more constrained, allowing only combinations of components with 0 or 1 weights (0 = exclusion, 1 = inclusion), providing the best subset of components. For the continuous weight optimization, in each group, composite weights were estimated via bootstrap resampling and cross-validation to find the set of weights that maximized the separation of the groups over the first 48 months of follow-up. Spline knots for models limited to 48-month follow-up were placed at 12, 24, and 36 months, post baseline. The median weight from this distribution for each outcome was used to form the composite to be evaluated for trial power. Details of each step are described in the following.

Longitudinal cognitive and functional measures were modeled using linear mixed-effects regression with a random intercept and slope and an unstructured covariance matrix for the random effects. Variance components were estimated conditional on conversion information. To capture departures from linearity in the trajectory of cognition and function, continuous time from the baseline test was parameterized using a 3-knot restricted cubic spline, with knots placed at 1, 3, and 5 years, post baseline. Differences in group trajectories were tested using linear mixed-effects regression with a random intercept and slope and an unstructured covariance matrix for the random effects. Variance components were estimated conditional on conversion information. In each training set, composite weights were estimated using bootstrap resampling and cross-validation to estimate power for a clinical trial based on participants who were both Aβ− and predicted to convert (pcCN), to make our results applicable to trials requiring Aβ+ for inclusion. Three-fold cross-validation was used for the pcCN analysis because of the reduced sample size.

The association between groups within each cohort and missing data was modeled using generalized mixed-effects regression with a binomial indicator for a missing visit. All analyses were done in R version 3.1.1 (The R Foundation for Statistical Computing, Vienna, Austria; www.r-project.org).

3. Results

3.1. Cohort characteristics

In the ADNI data set, 68 subjects converted to MCI during 7 years of follow-up while 70 subjects remained cognitively-normal throughout the same period. cCN subjects were older and had more APOE ε4 carriers compared to scCN (Table 1). There were no significant differences in gender or education. As described in the Section 2.5, we also identified a group of cognitively-normal subjects who were predicted to convert to MCI (pcCN, only including Aβ+ subjects). Characteristics of the pcCN group are shown in Supplementary Table 1.

One hundred thirty-seven Aβ+ subjects and 210 Aβ− subjects were included in the analysis. Aβ− subjects were older, less educated, and had more APOE ε4 carriers (Table 1). There was no difference in gender. A Kaplan-Meier plot showing the distribution of conversion times for the cCN, Aβ−, and Aβ+ groups is shown in Supplementary Fig. 2.
Of the 68 cCN participants, 56 had Aβ information: 31 (55.4%) were Aβ+ and 25 (44.6%) were Aβ−. Of the 70 sCN participants, 57 had Aβ information: 18 (31.6%) were Aβ+ and 39 (68.4%) were Aβ−.

### 3.2. Baseline cognitive/functional differences

When baseline cognitive/functional measures were compared in cCN versus sCN, cCN subjects performed worse on all 12 outcomes. Results with multiple comparison corrections are shown on the top left of Table 2. There were fewer differences on baseline cognitive/functional measures in Aβ+ versus Aβ− participants (top right of Table 2).

### 3.3. Longitudinal change

cCN subjects worsened significantly faster on 10 of the 12 cognitive and functional outcomes compared to sCN subjects, with the exception of the Boston Naming Test and Trails A over 7 years of follow-up (Fig. 1, Table 2, Supplementary Fig. 3). The largest effect size was in the CDR-SB, and the largest effect sizes among measures without functional items were in the iAVLT and the ADAS13. Longitudinal trajectories of the pcCN group are shown in Supplementary Fig. 4.

Aβ+ subjects worsened significantly faster on 6 of the 12 outcomes compared to Aβ− subjects (Fig. 1, Table 2). The largest effect size was in the ADAS13.

cCN subjects were more likely than sCN subjects to be missing data during the course of the 7 years of follow-up (log OR = 0.82, standard error = 0.15, p < 0.001). However, sCN subjects were selected to have a minimum follow-up time of 7 years. Aβ−-positivity was not associated with increased missingness (log OR = −0.04, standard error = 0.27, p = 0.87).

### 3.4. Composite weight distributions

The distributions from 1000 bootstrap samples of the composite weights that maximized the separation of the groups are shown in Supplementary Fig. 5. Composite weights were estimated separately for the 3 groups (cCN, pcCN, and Aβ+).

The largest contributing outcomes in the composite for cCN versus sCN were the 2 delayed memory recall measures (dMemory, dAVLT), CDR-SB, and the MMSE (top left of Supplementary Fig. 5). Outcomes with smaller, although nonzero, positive median weights, included Category Fluency, iMemory, Trails A, and the Boston Naming Test. When the functional measures were excluded, the delayed memory recall measures and MMSE remained the largest weighted outcomes and ADAS13 became more heavily weighted.

Composite weights that maximized the separation of pcCN and sCN subjects were also estimated. Using baseline information including demographics, APOE ε4 status, and cognitive/functional variables that were not heavily weighted in the true converter composite (ADAS13, Trails A & B, FAQ, Boston Naming Test, iAVLT, and iMemory), composite weights were estimated based on 32 pcCN participants. In reality, these 32 pcCN participants consisted of 25 converters and 7 nonconverters, resulting in a 78% positive predicted value from the model estimates. pcCN subjects were older, less educated, had more APOE-ε4 allele carriers, and had lower cognitive scores at baseline compared with sCN subjects, similar to cCN subjects (Supplementary Table 1). We then estimated composite weights for this cohort. These weights are shown in the middle row of Supplementary Fig. 5. Similar to the cCN composite, the main outcomes for the pcCN composite were dMemory, CDR-SB, and MMSE, but in contrast, included the Boston Naming Test. When functional measures were excluded, the ADAS13 carried more weight. Note that the pcCN were Aβ+ by design because we aimed to make our results applicable to a trial requiring Aβ+ for inclusion.

The composites for Aβ+ versus Aβ− were heavily weighted by ADAS13, FAQ, and MMSE (bottom left Supplementary Fig. 5). When functional measures were excluded, ADAS13 and MMSE dominated the composites.
3.5. Best subset components

The best subset results were similar to the continuous optimization results. For the cCN versus sCN comparison, 5 components provided the optimal cross-validated composite, with the MMSE, dMemory, dAVLT, CDR-SB, and Category Fluency selected in nearly all cross-validation folds. For the pcCN versus sCN comparison, 7 components were selected, including the MMSE, dMemory, dAVLT, CDR-SB, Category Fluency, and iMemory in nearly all folds and occasionally either ADAS13 or Trails A. For the Aβ+ versus Aβ− comparison, 3 components were selected—the MMSE, ADAS13, and FAQ. The power for these composites is described in the following.

3.6. Power

We estimated the power to detect a 30% slowing of decline using the average out-of-sample estimates of change and variance for each composite and group, over a range of sample sizes. The composite with flat weights across all measures was the best performing composite, attaining 80% power with 375 completers/arm in a hypothetical 30-month trial. Eighty percent power was attained with 450 completers per arm using the optimized cognitive/functional composite in a hypothetical 30-month clinical trial. Sixty-five percent of power was obtained with 500 completers per arm over a 30-month trial, using a
composite with cognition only. We also compared flat weight and optimized cognitive/functional composites in 48-month trials for Aβ+ pcCN subjects. They performed similarly (Supplementary Fig. 6).

The power estimates for the Aβ+ subjects are shown in the lower portion of Fig. 2. With similar sample sizes as the comparisons mentioned previously, power estimates never exceeded 40% with any type of composite.

3.7. Effect sizes and variance components

In Fig. 3, the magnitude of change, the within- and between-subject standard deviations, and effect sizes are plotted against the number of components included in the best subset composites, for cCN versus sCN and Aβ+ versus Aβ-. For both groups, the magnitude of change and both types of SD decrease with an increasing number of components included in the composites.

4. Discussion

The main findings of this study were as follows: (1) including participants with Aβ pathology as well as subtle cognitive dysfunction, predictive of progression to MCI, resulted in large gains in power estimates compared to participants with Aβ pathology alone; (2) further gains in power were obtained by including measures with functional items in the composite; (3) composites formed by applying equal weights to all 12 measures provided the highest estimates of cross-validated power, although similar to continuous weight optimization and best subset optimization; (4) as the number of components in the composite increased, the magnitude of change decreased, but both the within-subject and between-subject variance decreased, leading to an increase in effect size; (5) in cCN and pcCN participants, the composite measures selected via optimization were both delayed memory recall assessments, CDR-SB, MMSE, Category Fluency, and immediate memory recall; in Aβ+ participants, ADAS13, MMSE, and FAQ were...
selected, however, these composites did not outperform the equal weight composites when cross-validated in either group; and (6) only 55.4% of cCN subjects were \( A\beta^+ \), which explains part of the difference between our analysis of cCN and \( A\beta^+ \) subjects, and points to the importance of non-\( A\beta \)-mediated processes to explain development of cognitive and functional decline.

4.1. Power increase with predicted converters

Substantial increases in power estimates result when including pcCN subjects compared to \( A\beta^+ \) subjects, in all clinical trial simulations (Fig. 2). This might be expected when considering Fig. 1 and Table 2, where decline is limited both in magnitude and number of outcomes in the \( A\beta^+ \) subjects compared to cCN subjects, especially over the first 36 months. In contrast, the cCN subjects have already diverged from sCN subjects on several measures at baseline and continue to separate on delayed memory recall, global cognitive, and functional outcomes. Lower cognitive scores and continued decline in both the pcCN and cCN groups indicate that these participants are likely already in a later stage of disease at baseline compared with \( A\beta^+ \) participants. The lower power estimates using ADNI \( A\beta^+ \) subjects are consistent with estimates reported in the analysis of 2 \( A\beta^+ \) cohorts for the A4 composite (Donohue et al., 2014). Substantially shallower decline was observed in the cognitively-normal ADNI \( A\beta^+ \) subjects compared to the cognitively-normal \( A\beta^+ \) subjects from the Australian Imaging, Biomarkers & Lifestyle Flagship Study of Ageing (AIBL, Ellis et al., 2009; Donohue et al., 2014). The sharper decline seen in the \( A\beta^+ \) subjects in the AIBL cohort may be due to subtle cognitive dysfunction at baseline including a 0.4 point lower average MMSE score, 0.5 point lower delayed memory delayed recall score, as well as a 20% increase in \( APOE \varepsilon 4 \) allele carriers, compared with the ADNI \( A\beta^+ \) subjects. These subtle differences in baseline cognition and the increased proportion of \( APOE \varepsilon 4 \) carriers may account for the differences in power estimates, which are closer to the estimates of the pcCN cohort observed in this analysis. Taken together, these results point to the importance of assessing other baseline characteristics besides \( A\beta \) status when selecting preclinical populations.
for trial enrichment. This should come as no surprise given the vast literature on the variability of the clinical impact of Aβ pathology in elderly people, where a similar degree of Aβ pathology may be seen in people who are cognitively normal, slightly impaired or fully demented. This variability likely stems from individual differences in cognitive and brain reserve mechanisms, differences in the presence and spread of important copathologies (including tau pathology), and differences in the time that the individual has been exposed to Aβ pathology before testing. Additional sources of variation regarding the effect of Aβ pathology on cognition in cognitively-normal cohorts include biomarker modality (PET vs. CSF) and also choice of threshold for Aβ-positivity (Insel et al., 2014; Mattsson et al., 2014, 2015). The impact of this on the power of clinical trials, as found in our results, is in agreement with a previously proposed staging of preclinical AD (Sperling et al., 2011b), where subjects with a combination of positive Aβ biomarkers (including Aβ1-42 biomarkers) and subtle cognitive dysfunction are thought to be at much higher risk for future clinical deterioration compared to subjects with positive Aβ biomarkers alone (Vos et al., 2013).

4.2. Additional power increase with functional components

Including the CDR-SB and FAQ in either the optimized composite or the equal weight composite resulted in an additional increase in power over the cognitive composite in the pCN subjects, reaching 80% power with 375 completers per arm for a 30-month trial (Fig. 2). Including measures with functional items provided moderate improvements in power for the composites in Aβ+ subjects for trials less than 36 months, although power remained low. To convert from normal cognition to MCI, a subject must demonstrate a clinically meaningful level of functional decline. Steep decline is observed on both the CDR-SB (Fig. 1) and on the FAQ immediately after baseline in cCN subjects. Thus, it follows that including measures with functional assessments in a composite results in a more sensitive instrument, in a population of converters. However, because conversion status in not known at baseline, the inclusion of functional assessments in a prospective study will only improve sensitivity if information available at baseline can successfully identify subjects who are on the verge of functional decline. When functional measures are excluded, the weights for both the ADAS13 and the MMSE increase. This may reflect that poor scores on a global cognitive test are likely more correlated with functional decline compared to single domain measures. Aβ+ subjects do not show substantial decline on either CDR-SB or FAQ before month 48.

4.3. Functional and cognitive outcome selection

Delayed memory recall, the MMSE, and the CDR-SB were selected via optimization for both the cCN and the pCN composites. However, even the top-weighted measures had relatively low median weights, with 10 of the 12 measures having positive weights for cCN subjects, and 6 of 12 having positive weights for pCN subjects (Supplementary Fig. 5). The spread of the weights suggests that many domains are declining early in the conversion process. Thus, it follows that the equal weight composites performed well. The failure of the optimization to beat the equal weight composites suggests that using either continuous weights or best subset component selection results in overfitting the training sets and a subsequent reduction of test set power. Including a large number of components in a composite may smooth over aberrations in scores in a particular assessment from visit to visit within a subject, thus lowering the within-subject variance and improving signal to noise. Similarly, the equal weight composite provided the most power in Aβ+ participants, although power did not approach levels suitable for a phase III trial (Fig. 2).

4.4. Equal weight composite: effect size, magnitude of change, and variance

Reasons for slight increases in power with the equal weight composite become clear from inspection of Fig. 3. As the number of components included in the composite increases, the magnitude of change decreases. This would result in a decrease in effect size (if the variance is held constant) and subsequently, a decrease in power. If we start with the best single component and continue by adding additional components, the magnitude of change may become diluted as less-sensitive components are included in the composite. We might expect the effect size to drop with each additional component; however, both the within-subject and between-subject errors are decreasing at a rate that overcomes the decrease in the magnitude of change, resulting in an increasing effect size, as seen at the bottom of Fig. 3. The increase in effect size plateaus in the 6–10 component range for both the converter and Aβ+ groups. The decrease in within-subject variance is clear in both groups; however, the drop in between-subject variance is steeper for converters, likely due to more consistent decline across all components. Or alternatively, the converters’ scores are more variable, with more room for a reduction in within-subject variance when the number of composite components increases.

4.5. Outcome selection in other studies

The outcomes selected via optimization are consistent with the measures found to capture decline in other cohorts. The API composite in Presenilin 1 E280A mutation carriers includes the Word List Recall (CERAD), MMSE (orientation to time), and also Constructional Praxis and Raven’s Progressive Matrices (Ayutyanont et al., 2014). The API composite developed from normal to MCI or AD converters includes Category Fluency, Logical Memory II (dMemory), MMSE (orientation to time), and also Ravens Progressive Matrices Subset, and Symbol Digit Modalities (Langbaum et al., 2014). The A4 composite for Aβ-positivity includes the Total Recall score from the Free and Cued Selective Reminding Test, Logical Memory II (dMemory), MMSE, and the Digit Symbol Substitution Test (Donohue et al., 2014). Delayed memory recall, orientation, and processing speed are consistently selected domains.

A variety of approaches can be used to develop composites that are sensitive to change over time (Ard et al., 2015). The development of composite measures may require the comparison of a large number of combinations of items, especially if weights are considered, leading to an increased risk of overfitting and an inflated estimate of the sensitivity and statistical power of the constructed composite. A validation procedure in a sample outside that used to identify the items and weights will be critical to accurately assess the composite’s performance (Hendrix, 2012). As seen in our analysis, both types of optimization resulted in reduced power compared with the equal weight composites, likely due at least in part by overfitting the training sets. Importantly, the composites developed in this study and for the A4 study were evaluated out of sample. Neither study was able to reliably improve on equal weights.

4.6. Limitations

This study has several limitations. We evaluated assessments available in the ADNI neuropsychological battery, although it is possible or likely that there are other measures more sensitive to
decline in preclinical AD. We also did not consider item-level data from already formed composites, such as the orientation to time component of the MMSE (Langbaum et al., 2014), which may have affected the results due to carrying insensitive items along with more sensitive ones. We also make the assumption that a treatment will slow the progression of components selected for their fast decline. In reality, it is unknown which cognitive or functional components a treatment may affect and it is possible that an endpoint comprising slower progressing domains will yield more power. Additionally, we used restricted cubic splines to model the observed data and subsequently simulated clinical trials assuming an MMRM model. Maximizing the group trajectory differences assuming a spline model averages change over all time points to estimate the group curves, whereas the MMRM model allows change at each time point to be estimated more independently. Assuming an MMRM model for both steps of the analysis and allowing the weights to be differentially optimized according to trial length may yield different results. The ADNI cohort, with high levels of education possibly contributing to increased cognitive reserve, and also limited cognitive decline observed in the Aβ+ subjects compared with other cohorts, is potentially different from a population recruited for a clinical trial. The pcCN cohort is also considerably smaller with only 32 participants, reducing the stability of the estimates compared with the other cohorts. We used some of the same measures to predict conversion at screening and also track decline in the reference (equal weight) composite. It’s possible that a regression to the mean effect could result in a reduction of power. However, the equal weight composite remained the most reliably performing composite with considerable power.

5. Conclusion

Our results suggest preclinical AD subjects with lower cognitive scores at baseline decline more reliably across both cognitive and functional measures compared to Aβ+ subjects without signs of subtle cognitive dysfunction. This provides a challenge to designers of preclinical AD trials to identify the level of cognitive dysfunction to be required at screening that will result in further decline, allowing a treatment effect to be demonstrated. Later stage preclinical AD may represent a more feasible target population for clinical trials designed to slow cognitive decline. In this population, suitable power for a phase III trial can be achieved with considerably lower sample sizes while capturing both cognitive and functional change to demonstrate a clinically meaningful drug effect—both while initiating treatment in subjects who are still cognitively normal. Multiple measures of delayed memory recall, orientation, processing speed, as well as multiple functional measures should be considered when forming a composite. Finally, when selecting measures, erring on the side of too many components may be preferable to too few.

Disclosure statement

Mr. Insel, Dr. Mattsson, Dr. Hansson, and Dr. Mackin report no disclosures. Dr. Donohue was a consultant for Bristol-Myers Squibb. Dr. Aisen serves on a scientific advisory board for NeuroPhage; has served as a consultant to Elan, Wyeth, Eisai, Schering-Plough, Bristol-Myers Squibb, Eli Lilly and Company, NeuroPhage, Merck, Roche, Amgen, Genentech, Abbott, Pfizer, Novartis, Bayer, Astellas, Dainippon, Biomarin, Solvay, Otsuka, Daiichi, AstraZeneca, Janssen, Medication, Ichor, Toyama, Lundbeck, Biogen Idec, IPerian, ProBio-drug, Somaxon, Biotie, Cardues, Anavex, Kyowa Hakko Kirin Pharma, and Medtronic; and receives research support from Eli Lilly and Baxter and the NIH [NIA U01-AG10483 [PI], NIA U01-AG024904 [Coordinating Center Director], NIA R01-AG030048 [PI], and R01-AG16381 [Co-PI]]. Dr. Weiner has been on scientific advisory boards for Pfizer and BOLT International; has been consultant for Pfizer Inc.; Janssen, KL Associates, Easton Associates, Harvard University, inThought, INC Research, Inc, University of California, Los Angeles, Alzheimer’s Drug Discovery Foundation, and Sanofi-Aventis Groupe; has received funding for travel from Pfizer, ADPD meeting, Paul Sabatier University, Novartis, Tohoku University, MGI Group, France, Travel eDreams, Inc, Neuroscience School of Advanced Studies (NSAS), Danone Trading, BV, CTAD ANT Congress; serves as an associated editor of Alzheimer’s & Dementia; has received honoraria from Pfizer, Tohoku University, and Danone Trading BV; has research support from Merck, Avid, DOD, and VA; and has stock options in Synarc and Elan.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.neurobiolaging.2016.08.017.

References

A R T I C L E   I N F O

Keywords:
Alzheimer’s disease  
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Cognition

A B S T R A C T

Objective: Estimate the time-course of the spread of key pathological markers and the onset of cognitive dysfunction in Alzheimer’s disease.

Methods: In a cohort of 335 older adults, ranging in cognitive functioning, we estimated the time of initial changes of Aβ, tau, and decreases in cognition with respect to the time of Aβ-positivity.

Results: Small effect sizes of change in CSF Aβ/42 and regional Aβ PET were estimated to occur several decades before Aβ-positivity. Increases in CSF tau occurred 7–8 years before Aβ-positivity. Temporoparietal tau PET showed increases 4–5 years before Aβ-positivity. Subtle cognitive dysfunction was observed 4–6 years before Aβ-positivity.

Conclusions: Increases in tau and cognitive dysfunction occur years before commonly used thresholds for Aβ-positivity. Explicit estimates of the time of these events provide a clearer picture of the time-course of the amyloid cascade and identify potential windows for specific treatments.

Introduction

Disconcerting clinical trial results for the treatment of Alzheimer’s disease (AD) have led to a shift toward earlier intervention, focusing on the early clinical or presymptomatic phases, when biomarkers are needed to identify the disease. The amyloid cascade (Hardy and Selkoe, 2002) is thought to start with elevated levels of two key amyloids in the brain, β-amyloid (Aβ) and tau, and end with severe cognitive and functional impairment (Jack et al., 2010). Growing evidence suggests that an early sign that the cascade has begun is change in cerebrospinal fluid (CSF) Aβ, potentially detectable prior to significant Aβ deposition in the brain as measured by positron emission tomography (PET) (Palmqvist et al., 2016). This accumulation of Aβ has been suggested to be followed by increases in CSF tau and the spread of tau pathology beyond the temporal lobe (Braak and Braak, 1991; Schöll et al., 2016). The build-up and spread of these two brain pathologies is paralleled by gradual cognitive and functional decline (Zetterberg and Mattsson, 2014).

Previous neuropathological and biomarker data suggest that the overall time-course of AD is several decades (Li et al., 2017; Villedemagne et al., 2013). In autosomal dominant AD, the estimated years to clinical onset has been used to estimate the time-course of different biomarkers in AD (Bateman et al., 2012). However, the time-course of the spread of Aβ and tau and the onset of clinical symptoms in sporadic AD is unknown. With repeated measures of Aβ over time, the level and rate of change with respect to the key initiating AD pathology may offer a measure of disease progression in sporadic AD. The duration of amyloid positivity (chronicity) has been shown to be associated with increased tau pathology and faster cognitive decline and valuable in explaining heterogeneity in early disease progression (Koscik et al., 2020). With level and change information, the time from the threshold for significant Aβ pathology can be estimated within individuals, providing the temporal disease progression information important for evaluating biomarker trajectories. Without longitudinal information, cross-sectional studies frequently categorize subjects into two groups – those below a threshold for significant pathology and those above, where subjects just below the threshold who will cross over within months are considered pathologically equivalent to subjects who will not cross over for decades. By incorporating longitudinal information, disease progression with respect to Aβ pathology can be represented to reflect its continuous nature, resulting in a more...
powerful way to model the relationship between $A\beta$ and downstream processes.

The aims of this study were to demonstrate the utility and predictive ability of the time-from-$A\beta$-positivity (TFAP+$\beta$) formulation and to evaluate the relationships between TFAP+$\beta$- and downstream biomarker and cognitive responses in order to estimate the time of the earliest signs of progression in sporadic AD. Using serial $^{18}$F-florbetapir ($A\beta$) PET measurements, rates of change of $A\beta$ were estimated and used to calculate the time-from-threshold for each subject. These subject-specific estimates of the proximity to the threshold for $A\beta$-positivity ($A\beta(+)$) were then used to model the trajectories and temporal ordering of other key markers in AD including CSF $A\beta$42, regional $A\beta$ PET, several measures of tau including CSF phosphorylated (P-tau) and total tau (T-tau), regional $^{18}$F-flortaucipir (AV-1451) tau PET, and cognition. Estimates of the time and ordering of these pathophysiological changes may facilitate the design of future prevention trials and identify a window for early treatment.

Materials and methods

Standard protocol approvals, registrations, and patient consents

This study was approved by the Institutional Review Boards of all of the participating institutions. Informed written consent was obtained from all participants at each site.

Data availability

All data is publicly available (http://adni.loni.usc.edu/). R code is available on Github.

Participants

Data were obtained from the Alzheimer’s Disease Neuroimaging Initiative (ADNI) database (http://adni.loni.usc.edu/, www.adni-info.org) on 1/21/2020. An initial analysis was done on ADNI participants with available $A\beta$ PET data (In N = 962 $A\beta$-cognitively unimpaired (CU), $A\beta+$-cognitively unimpaired (CU+), $A\beta+$ MCI and $A\beta+$ AD), to facilitate the estimation of TFAP+$\beta$, though not all 962 were included in the analysis of the main outcomes. Participants were classified as cognitively unimpaired if they had an MMSE score of 24–30, CDR score of 0, no memory complaint, and a Logical Memory II subscale of the Wechsler Memory Scale-Revised score ≥ 9 for 16 years of education, ≥ 5 for 8–15 years of education, and ≥ 3 for 0–7 years of education. Participants were classified as MCI if they had an MMSE score of 24–30, a CDR score score of 0.5 as well as a memory box score of 0.5, and a Logical Memory II score ≤ 8 for 16 years of education, ≤ 4 for 8–15 years of education, and ≤ 2 for 0–7 years of education. Subjects were classified as having AD dementia if they had a memory complaint, met the same criteria for Logical Memory as the MCI group, had a CDR score of 0.5 or 1, and met the National Institute of Neurological and Communicative Disorders and Stroke–Alzheimer’s Disease and Related Disorders Association criteria for probable AD. The population in the primary analyses of PET and cognitive outcomes only included ADNI participants with available measurements of $A\beta$ and tau PET and cognition. Of these, all cognitively unimpaired (CU- and CU+), prodromal AD ($A\beta+$ MCI) and $A\beta+$ AD dementia participants were included in the analysis, where $A\beta$-positivity was defined using a previously established threshold (Standardized Uptake Value Ratio, SUVR = 1.10) (Joshi et al., 2012). $A\beta$- MCI (N = 224, including $A\beta$- CU to MCI progressed) and $A\beta+$ “AD dementia” subjects (N = 51, including $A\beta+$ MCI to AD dementia progressed; we consider these to be misdiagnosed, because we assume AD requires $A\beta(+)$ were not included in the main analysis given our aim to model head to head comparisons of initial biomarker and cognitive changes of individuals on the AD continuum and not other diseases. In order to estimate the time of emerging cognitive decline associated with increasing $A\beta$, those with cognitive impairment, but low levels of $A\beta$ were excluded. MCI and AD participants were considered $A\beta+$ if their SUVR remained below the threshold at all scans during follow-up. Visualizations of their biomarker data are included for comparison in Figs. 2–4 (see Figure legends). Additional description is included in the statistical analysis section.

Cerebrospinal fluid biomarker concentrations

Cerebrospinal fluid (CSF) samples were collected at baseline by lumbar puncture in a subsample (N = 185). CSF $A\beta$42, total tau (T-tau) and phosphorylated tau (P-tau) were measured by an xMAP assay (NINCA AlzBio3, Ghent, Belgium, Fujirebio), as described previously (Olsson et al., 2005; Shaw et al., 2009).

PET imaging

Methods to acquire and process $A\beta$ (18F-florbetapir) PET image data were described previously (Jagust et al., 2015; LANDAU et al., 2012). Briefly, florbetapir image data were acquired 50–70 min postinjection, and images were averaged, spatially aligned, interpolated to a standard voxel size, and smoothed to a common resolution of 8 mm full width at half maximum. We used an a priori defined threshold for $A\beta$-positivity (SUVR=1.1) (ADNI, 2012; JOSHI et al., 2012) applied to the ratio of the average of the four target regions (temporal, cingulate, frontal, and parietal lobes) and the cerebellum, in the estimation of time-from-$A\beta$-positivity, described in detail below. In a second part of the analysis, five $A\beta$ PET ROI outcomes were considered (LANDAU and JAGUST, 2015; MORMINO et al., 2009), (1) the temporal lobe (middle and superior temporal lobe), (2) the parietal lobe (precuneus, supramarginal, inferior and superior parietal lobe), (3) the cingulate gyrus (isthmus, posterior, caudal and rostral anterior cingulate), (4) the frontal lobe (pars opercularis, pars triangularis, pars orbitofrontalis, caudal/rostral middle frontal, medial/lateral orbitofrontal, frontal pole, and superior frontal lobe), and (5) a composite of regions thought to be early in accumulating $A\beta$ (precuneus and posterior cingulate) (Palmequist et al., 2017). These ROIs comprise the regions included in the global composite, grouped into individual lobes plus an additional early ROI. 18F-florbetapir ROIs were expressed as SUVRs with a cerebellar reference region.

Methods to acquire and process tau (18F-flortaucipir) PET image data were described previously (MASS et al., 2017). Six tau ROI outcomes, corrected for partial-volume, were considered: (1) the medial temporal lobe (MTL) (amygdala, entorhinal and parahippocampal cortex; from Braak stage I and II), (2) the lateral temporal lobe (LTL) (inferior/middle/superior temporal lobe, banks of the superior temporal sulcus, transverse temporal lobe, temporal pole; from Braak stage IV and V), (3) the medial parietal lobe (MPL) (isthmus cingulate, precuneus; from Braak stage IV and V), (4) the lateral parietal lobe (LPL) (inferior/superior parietal lobe, supramarginal; from Braak stage V), (5) frontal lobe (pars, orbitofrontal and middle/superior frontal lobe; from Braak stage V), and (6) the occipital lobe (cuneus, lingual, pericalcarine, and lateral occipital lobe; from Braak stage III, V, and VI). 18F-flortaucipir ROIs were expressed as SUVRs with an inferior cerebellar gray matter reference region. Scanner type and site were evaluated for their association with PET outcomes through covariate adjustment. Full details of PET acquisition and analysis can be found at http://adni.loni.usc.edu/methods/.

Cognition

Cognitive measures assessed included the Mini-Mental State Examination (MMSE) as a measure of global cognition, and the Preclinical Alzheimer’s Cognitive Composite (PACC), as a measure of early AD-related cognitive changes. The PACC comprised the MMSE, the Logical Memory Delayed Word Recall from the Wechsler Memory Scale, the Alzheimer’s Disease Assessment Scale—Cognitive Subscale Delayed Word Recall, and the Trail Making Test part B (reversed such
that high scores indicated better performance and log transformed) (Donohue et al., 2017, 2014). The PACC was constructed from available data in the sample. Components were z-transformed, summed and scaled to the baseline scores of the Aβ+ CU.

**Statistical analysis**

The aims of these analyses were to (1) demonstrate the utility and predictive ability of the TFA+ formulation of Aβ information and (2) evaluate the relationship between TFA+ and CSF, PET, and cognitive responses in order to estimate the time of the earliest signs of progression. The overall analysis consisted of two steps. Step one was estimating TFA+ based on the longitudinal measures of global Aβ PET SUVR. In step two, TFA+ estimates were used to predict cross-sectional measures of regional tau and Aβ PET, CSF and cognitive outcomes. To demonstrate the value of the TFA+ measure, we did head-to-head comparisons of (i) TFA+ vs (ii) intercepts and slopes of longitudinal Aβ PET, modeled separately, to predict the outcomes. These comparisons are described in detail below.

Because TFA+ was not directly observed, in step one, linear mixed-effects models were fit to all available longitudinal global Aβ PET SUVR data to estimate subject-specific intercepts and slopes of Aβ pathology.

For the ith individual at the jth measurement time, $y_{ij} = \beta_0 + \beta_1 t_j + \beta_2 b_j + \epsilon_{ij}$, where $y_{ij}$ is Aβ SUVR, $\beta_0$ and $\beta_1$ are the fixed intercept and slope over time, $t_j$ is time (years from baseline), $b_j$ and $h_j$ are the random intercept and slope over time, and $\epsilon_{ij}$ is a Gaussian distribution error term.

Because Aβ slopes are unlikely to remain constant over long periods of time as subjects move toward and away from the Aβ threshold, natural splines (Hastie and Tibshirani, 1990) were used to estimate the nonlinear shape of the slopes with respect to baseline Aβ, using quantile regression (Koenker and D’Orey, 1987). Rather than modeling the mean Aβ slope with respect to baseline Aβ, quantile regression provides a separate curve for each quantile, allowing the relationship between slope and intercept to differ depending on the location in the distribution of Aβ slope.

For a random variable $X$, with distribution function $F$, the rth quantile of $X$ is defined as, $Q_r(X) = F^{-1}(r)$. Taking the sum of the random variables $X_1 + X_2 + h_1$, gives the subject-specific estimates of the slope over time of Aβ SUVR. Similarly, taking the sum, $t_j = b_0 + b_0$, gives subject-specific estimates of the intercept of Aβ SUVR. Quantile curves were estimated by regressing $S$ on $t$ for $r \in (0, 1)$,

$Q_r(t) = \beta_0 + \beta_1 t + \beta_2 b + \epsilon_r$,

where $X_1, \ldots, X_2$ is the k-dimensional natural spline basis for $t$. The curve shape was selected by AIC.

For each subject, TFA+ was estimated by integrating over each subject’s quantile curve from the subject’s intercept to the threshold for Aβ+ positivity (PET SUVR = 1.1). For example, for a subject with a baseline SUVR of 1.2 and a slope in the 0.6 quantile, TFA+ was taken to be the time it would take to go from SUVR = 1.1 to 1.2, using the slope estimates from the quantile curve. For incremental changes on the x-axis (baseline SUVR), the time required to travel the incremental distance is equal to distance/rate. Using the trapezoidal rule (Atkinson, 1989), TFA+ is the sum of these incremental times spanning SUVR = 1.1–1.2. An example of calculating TFA+ is given in the top left panel of Fig. 1. Sensitivity analyses were done to determine the effect of varying the threshold for Aβ+.

We repeated the estimation of TFA+ using an early threshold (SUVR 1.07) and a late threshold (SUVR 1.13).

To evaluate the accuracy of the TFA+ estimates, we compared the observed times of Aβ+ to the estimated times of Aβ+ values in participants who were Aβ+ at baseline and became Aβ+ during follow-up. Observed time of Aβ+ occurred in the interval between the last Aβ− scan and the first Aβ+ scan. The observed time was calculated as a weighted average of the two scan times, weighted proportionally to the scan where the participant was closest to hitting the threshold. Observed and estimated values were compared in $N = 37$ participants who crossed the threshold for Aβ+ and remained Aβ+ throughout follow-up. We also compared observed and estimated values in 44 participants including the original 37 plus seven additional subjects who crossed the threshold but had a subsequent negative scan.

Our analyses aim to model participants who are ostensibly on the AD trajectory and had calculable TFA+, i.e., they must be Aβ+ accumulators (positive rates of accumulation). Therefore, of the 982 participants with Aβ PET, we excluded $N = 20$ participants with negative Aβ+ accumulation rates (negative rates were largely driven by one early high Aβ+ PET measure), we also excluded $N = 6$ participants with low levels of Aβ+ and accumulation rates such that they were predicted to become Aβ+ later than 120 years of age (biomarker data from these subjects are included for visual comparisons in Figs. 3–5, see Figure legends). Fig. 1 shows the flow of participant inclusion. We included subjects where the TFA+ metric indicated very early accumulation of Aβ, but for participants estimated to have become Aβ+ before age 40 ($N = 25$, median estimated age at Aβ+ = 30, IQR: 27 to 34), we truncated TFA+ to age 40, based on previously described rates of Aβ+positivity in middle age (Jansen et al., 2015). These were mostly MCI and AD participants, APOE ε4 carriers, in their mid 60 s to late 70 s.

In the second step, the relationship between TFA+ and the responses was modeled using monotone penalized regression splines. The model takes the form,

$y_i = f(TFA+, \tau) + \epsilon_i = a_1(TFA+, \tau) + \ldots + a_q(TFA+, \tau) + \epsilon_i$,

where $y_i$ is one of the PET, CSF, or cognitive responses, measured cross-sectionally, and $f$ is a smooth monotone function, represented by $a_1, \ldots, a_q$ basis functions. Generalized cross-validation was used to control the basis dimension $q$ and the degree of smoothing (Wood, 1994).

Cognitive responses were covaried for age, gender and education; CSF Aβ42, T-tau, P-tau and PET measures were covaried for age and gender.

Because the variance of the outcomes increases with advancing pathology and several outcomes contained clusters of extreme values, resulting in large residuals, we repeated step two of the analyses using M-estimation to provide robust estimates with robust standard errors (Huber and Ronchetti, 1981). This model takes the same form as described above, but is fit using iteratively reweighted least squares in order to downweight outliers.

In order to account for the variance across steps 1 and 2, the entire process was repeated in 500 bootstrap samples to estimate 95% confidence intervals for the association between TFA+ and the responses.

To assess the predictive ability of TFA+, we compared (i) models using TFA+ vs (ii) models using both the subject-specific intercepts and slopes of longitudinal Aβ, to predict the responses. Two models for each response were compared. In model 1, responses were regressed on TFA+ using penalized regression splines as described above, adjusting for covariates. In model 2, responses were regressed on both Aβ+ intercepts and slopes using penalized regression splines, adjusting for covariates. Model 1 and 2 were compared using the Akaike Information Criterion (AIC) (Akaike, 1974).

Meaningful effect sizes of change in increase in pathology or decrease in cognition with respect to TFA+ were estimated as part of step two of the analysis. A Cohen’s d effect size of 0.2 SD was considered small, 0.5 SD was considered medium, and a 0.8 SD effect was considered large (Cohen, 1988). A 0.2 standard deviation (SD) change from the mean response at the longest times (least pathological) from Aβ+positivity was taken to be the initial point of meaningful change. A 0.5 SD change was also shown as a more substantial effect size of change. For example, if the estimated mean PACC score at the lowest level of Aβ was 0 and the estimated mean PACC score at the time of Aβ+positivity was −0.5 and the residual SD of the PACC was 1.5, then the effect size at the time of Aβ+positivity = (−0.5 − 0)/1.5 = −0.33. A drop of 0.33 points on the PACC would be considered between a small (0.2) and medium (0.5) effect size, according to Cohen’s guidelines for interpreting the magnitude of the effects.
effect sizes. We also estimated change, 95% confidence intervals with bootstrap-estimated 2.5 and 97.5 quantiles, and statistical significance of change for each response at TFA\(\beta^+ = 0\), the time of \(\beta^+\)-positivity, with bootstrap-estimated standard errors.

Associations between missing data (CSF subsample vs full cohort) and demographics and TFA\(\beta^+\) were evaluated using logistic regression with a binomial indicator for missing data. Baseline associations between demographics and TFA\(\beta^+\) were assessed using Pearson correlation for age and years of education and a t-test for gender. Associations between diagnosis and demographics were assessed using F and t-tests for continuous variables and \(\chi^2\) tests for categorical variables. P-values were 2-sided and considered significant for \(p < 0.05\). A drop of 2 or more in AIC was considered meaningful model improvement. All analyses were done in R v4.0.0 (www.r-project.org).

**Results**

**Cohort characteristics**

Two-hundred and twenty-seven CU (127 \(\beta^-\) and 100 \(\beta^+\)) and 70 \(\beta^+\) MCI and 38 AD participants were included in the analysis. The diagnostic groups varied by mean age, sex, years of education, and proportion of APOE \(\epsilon 4\) (Table 1). The CU- group was significantly younger than all other diagnostic groups (\(p < 0.04\)). The MCI group had a significantly smaller proportion of females than both the CU- group (\(p = 0.02\)) and the CU+ group (\(p = 0.05\)). The MCI group had significantly lower mean years of education compared to the CU- group (\(p = 0.04\)) and the CU+ group (\(p = 0.02\)). The AD group also had significantly lower mean years of education compared to the CU- group (\(p = 0.01\)) and the CU+ group (\(p = 0.005\)). The CU- group had a significantly smaller proportion of APOE \(\epsilon 4\) carriers than all other diagnostic groups (\(p < 0.003\)).

**A\(\beta\) PET and estimation of TFA\(\beta^+\)**

TFA\(\beta^+\) was estimated with a median of 3 (range: 1 to 5) A\(\beta\) PET scans per participant. The average time between first and last scan was 3.3 years (SD=2.9) and the average time between scans was 2.2 years (SD=0.8). PET data came from 17 types of scanners across 58 sites. Neither site nor scanner type were associated with any A\(\beta\) PET outcome (\(\Delta\)AIC > 12) or tau PET outcomes (\(\Delta\)AIC > 19) and were not included in subsequent models. The correlation between subject-specific random intercepts and slopes was 0.32 (0.06 to 0.55). Residuals from the mixed model of repeated global A\(\beta\) PET SUVRs appeared approximately normal and with constant variance over time.
Across diagnoses, TFAβ+ ranged from −29 to 46 years, where higher (positive) TFAβ+ values indicate more time spent with a significant Aβ burden. The CU- group had a significantly lower mean TFAβ+ compared to all other diagnostic groups (p<0.001). The CU+ group had a significantly lower mean TFAβ+ compared to the MCI group (p<0.001) and the AD group (p<0.001), and the MCI group was significantly lower than the AD group (p=0.02).

Higher TFAβ+ was significantly associated with older age (r=0.28, 95% CI: 0.17 to 0.37, p<0.001), lower education (r=−0.15, 95% CI: −0.25 to −0.04, p=0.01) and APOE ε4-positivity (mean TFAβ+ in APOE ε4= 3.3 (SD=16.1) years and mean TFAβ+ in APOE ε4+ = 13.6 (SD=15.8) years, p<0.001). TFAβ+ was not associated with sex (mean TFAβ+ = 9.4 (SD=17.1) and 6.8 (SD=16.4) in males and females, respectively, p=0.16). Within-diagnosis TFAβ+ distributions are shown on the bottom right panel of Fig. 2. Quantile curves of the relationship between Aβ intercepts and slopes are also shown in the top right panel of Fig. 2, displaying the variation of acceleration of Aβ deposition over different levels of baseline Aβ.

TFAβ+ estimates were not sensitive to alternative thresholds for Aβ+ beyond a shift reflecting an earlier or later threshold. When the earlier threshold (SUVR 1.07) was used rather than SUVR 1.10, TFAβ+ estimates were shifted a median of 3.2 years earlier, but remained almost perfectly correlated with TFAβ+ using the SUVR 1.10 threshold (r=0.996). Similarly, when the late threshold was used (SUVR 1.13),
TFAβ⁺ estimates shift a median of 3.1 years later, but also remained almost perfectly correlated with TFAβ⁺+ using the SUVR 1.10 threshold (p=0.997).

**TFAβ⁺ performance**

TFAβ⁺ was highly correlated with observed time of Aβ⁺ (r=0.93, 95% CI: 0.87 to 0.97, p<0.001, bottom left panel of Fig. 2). When also including the seven participants with a subsequent negative scan after their initial positive scan, the correlation between TFAβ⁺+ and the observed time of Aβ⁺ was 0.95, 95% CI: 0.80 to 0.94, p<0.001.

When comparing the performance of TFAβ⁺+ versus using Aβ intercepts and slopes as separate predictors, TFAβ⁺+ significantly outperformed separate intercepts and slopes most, but not all of the time. TFAβ⁺+ significantly outperformed covariate only models for all outcomes. Using TFAβ⁺ resulted in significantly better prediction of MTL tau (AIC_{TFAβ⁺}=345.1, AIC_{IntSlope}=360.2, AIC_{IntSlopeAIC}=484.8), MPL tau (AIC_{TFAβ⁺}=467.9, AIC_{IntSlope}=472.4, AIC_{IntSlopeAIC}=532.9), occipital lobe tau (AIC_{TFAβ⁺}=272.1, AIC_{IntSlope}=274.2, AIC_{IntSlopeAIC}=325.8), CSF Aβ (AIC_{TFAβ⁺}=1825.1, AIC_{IntSlope}=1827.4, AIC_{IntSlopeAIC}=1902.1), CSF T-tau (AIC_{TFAβ⁺}=1854.4, AIC_{IntSlope}=1863.4, AIC_{IntSlopeAIC}=1902.9), MMSE (AIC_{TFAβ⁺}=1534.8, AIC_{IntSlope}=1549.4, AIC_{IntSlopeAIC}=1600.4), and the PACC (AIC_{TFAβ⁺}=1251.2, AIC_{IntSlope}=1261.8, AIC_{IntSlopeAIC}=1347.4). There was no difference between TFAβ⁺+ and separate intercepts and slopes for LTL tau (AIC_{TFAβ⁺}=398.6, AIC_{IntSlope}=399.5, AIC_{IntSlopeAIC}=471.5) and CSF P-tau (AIC_{TFAβ⁺}=1711.3, AIC_{IntSlope}=1709.6, AIC_{IntSlopeAIC}=1748.3) and separate intercepts and slopes was significantly better than TFAβ⁺ in predicting frontal lobe tau (AIC_{TFAβ⁺}=194.2, AIC_{IntSlope}=188.4, AIC_{IntSlopeAIC}=245.9) and LPL tau (AIC_{TFAβ⁺}=444.9, AIC_{IntSlope}=441.9, AIC_{IntSlopeAIC}=512.3).

**Regional Aβ PET**

Five regional ROIs (precuneus + posterior cingulate, frontal lobe, cingulate gyrus, temporal and parietal lobes) are shown plotted against TFAβ⁺ in Fig. 3. All 5 regions were estimated to reach a small, but meaningful (0.2 SD) increase in SUVR between 12 and 15 years before Aβ⁺-positivity, i.e. TFAβ⁺ = 0. Effect sizes over the span of TFAβ⁺ are shown in Fig. 3. At TFAβ⁺ = 0, all regions showed large, significant increases in SUVR (ΔSUVR ≥ 0.11, p ≤ 0.01) with the precuneus + posterior cingulate composite showing the largest increase (ΔSUVR = 0.16, p < 0.01) and the temporal lobe showing the smallest (ΔSUVR = 0.11, p < 0.01). Effect sizes for all regions were large (≥1) by the time of Aβ⁺-positivity. Table 2 summarizes the values of the outcomes at the longest times before Aβ⁺, i.e. the least pathological TFAβ⁺-time. Table 2 also shows the value and change of each outcome at the time of Aβ⁺-positivity (TFAβ⁺ = 0), p-value, the effect size of change of each outcome, the 0.2 SD change point with respect to TFAβ⁺, and corresponding 95% confidence intervals.

Analyses of regional Aβ PET outcomes were repeated using robust regression with robust standard errors. The dashed blue curves in Fig. 3 depict the robust fit. The robust curves are similar to the unweighted regression curves with some mild flattening in the TFAβ⁺ = 5 to 25 year range. The 0.2 SD change point estimates for the increase in SUVR ranged from 18 to 20 years before Aβ⁺-positivity (compared to 12–15 years before Aβ⁺-positivity in the main analyses). Similar to the unweighted analyses, all regions showed significance of Aβ at TFAβ⁺ = 0 (p<0.01).

**CSF**

CSF responses are plotted against TFAβ⁺ in Fig. 4. A 0.2 SD drop in CSF Aβ42 was estimated to occur 29 years before Aβ⁺-positivity (TFAβ⁺ = −29). At TFAβ⁺ = 0, CSF Aβ42 showed a very large effect size (ΔAβ42 = −68 ng/L, p<0.01, effect size = −1.99). At TFAβ⁺ = −2, or two years before Aβ⁺-positivity, the population curve passes through a previously published CSF Aβ42 threshold for Aβ-positivity (192 ng/L) (Shaw et al., 2009).

A 0.2 SD increase in CSF T-tau and P-tau was estimated to occur 7–8 years before the time of Aβ⁺-positivity (TFAβ⁺ = −7 and −8, respectively). At TFAβ⁺ = 0, significant increases of medium effect size of T-tau (ΔT-tau = 19 ng/L, p = 0.04, effect size = 0.46) and P-tau (ΔP-tau = 12 ng/L, p = 0.04, effect size = 0.47) were observed.

Robust curves are shown in dashed blue in Fig. 4. The change point estimate was 26 years before Aβ⁺-positivity for the decrease in CSF Aβ, 13 years before Aβ⁺-positivity for CSF P-tau, and 8 years before Aβ⁺-positivity for CSF T-tau. A more substantial flattening of the curves can be seen in both CSF P-tau and T-tau for TFAβ⁺ > 0. The effect size for CSF T-tau at TFAβ⁺ = 0 remained almost identical (0.47, p = 0.03) and the effect size for CSF P-tau increased moderately to 0.56 (p = 0.01). The effect size for CSF Aβ42 increased to −2.52 at TFAβ⁺ = 0 and remained significant (p<0.01).

In comparing the CSF subsample (N = 185) to the full cohort, missing CSF Aβ42 (or P-tau) was not associated with age (OR=0.996, p = 0.42), sex (OR=1.08, p = 0.15), or TFAβ⁺ (OR=1.00, p = 0.96). Missing CSF T-tau was not associated with age (OR=0.996, p = 0.32), sex (OR=1.07, p = 0.22), or TFAβ⁺ (OR=1.00, p = 0.72).

**Tau PET**

Six regional ROIs (MTL, LTL, MPL, LPL, frontal and occipital lobes) are shown plotted against TFAβ⁺ in Fig. 5. Five of the six regions were estimated to reach a 0.2 SD increase in SUVR 3–5 years before Aβ⁺-positivity, with the occipital lobe reaching a 0.2 SD increase at the time of Aβ⁺-positivity. Effect sizes over the span of TFAβ⁺ are shown in Fig. 5. At TFAβ⁺ = 0, four regions (MTL, LTL, MPL, LPL) showed significant increases in SUVR (ΔSUVR > 0.14, p ≤ 0.03) with the MTL showing the largest effect size (0.36). The frontal and occipital lobes did not increase significantly by TFAβ⁺ = 0 (ΔSUVR = 0.09 (p = 0.06) and 0.07 (p = 0.13), respectively). Estimates are summarized in Table 2.

**Cognition**

Cognitive measures are shown in Fig. 6. The MMSE showed a 0.2 SD drop six years before Aβ⁺-positivity, followed by the PACC four years before Aβ⁺-positivity. Neither measure decreased significantly by the time of Aβ⁺-positivity (ΔMMSE = −0.71, p = 0.13, effect size=−0.30; ΔPACC = −0.50, p = 0.10, effect size=−0.32).

Robust curves are shown in dashed blue in Fig. 6 and show mild flattening for TFAβ⁺ > 0, compared to the unweighted analyses. The change point estimates for the decrease in cognitive scores was two years before Aβ⁺-positivity for MMSE and four years before Aβ⁺-positivity for the PACC. The robust estimate for the effect size of decrease in MMSE scores was reduced to −0.23 but became statistically significant (p = 0.03). The robust estimate for the effect size of decrease in PACC scores was similar (−0.30), and also became statistically significant (p = 0.03).

Summary curves and 0.2 SD change points for some of the earliest changing measures of each outcome type (CSF Aβ and P-tau, precuneus + posterior cingulate Aβ PET, MTL tau PET and the PACC) are shown in Fig. 7.

**Discussion**

Several biological processes develop over time in sporadic AD, including accumulation of Aβ and tau across wide areas of the brain, as well as cognitive decline. Based on the amyloid cascade hypothesis, a relevant overarching time scale of the disease processes could be based
on the development of Aβ pathology (Kocskí et al., 2020). Integrating Aβ PET level and rate of change information places each individual on a pathological timeline. While this timeline, represented in these analyses by TFAβ+, was more closely associated with tau PET, CSF and measures of cognition in most measures, compared with intercept and slope information modeled separately, the main advantage is that it is parameterized directly to estimate the time of downstream events in the amyloid cascade. We estimated several major milestone events of AD progression including a small drop in CSF Aβ42 29 years before Aβ-positivity and a small increase in regional Aβ PET deposition 15 years before Aβ-positivity. Using the biomarkers tested here, the first changes in CSF Aβ42 may define the onset of AD. Small increases in tau pathology were estimated to occur 7–8 years before Aβ-positivity, as measured by CSF and 5 years before, as measured by PET. More substantial and statistically significant increases in CSF as well as temporoparietal tau PET were detected by the time of Aβ-positivity. Small effects of cognitive dysfunction occurred 4–6 years before Aβ-positivity, coinciding with previous reports (Insel et al., 2017). These findings provide a
general time scale for initial changes in sporadic AD, which may inform clinical trials aimed at specific stages of the disease.

Once beyond the threshold for Aβ-positivity, there is a substantial increase in the variance of the tau and cognitive responses. A handful of participants show large increases, especially in tau PET, and large decreases in cognition, resulting in clusters of outliers. These outliers appear to have marked influence on both the shape of the curves and the estimates of the variance, as shown by the difference between the unweighted and the robust analyses. The robust curves are generally flatter beyond the threshold for Aβ-positivity, less influenced by extreme values. The curves are reasonably similar prior to Aβ-positivity, although the overall variance estimates are smaller, resulting in earlier estimates of change points for several of the outcomes and more significant differences at the threshold for Aβ-positivity. In both sets of analyses, significant increases in both CSF tau and tau PET are observed to occur by the time of Aβ-positivity.

A 0.2 SD difference, a small, but meaningful increase in levels of CSF tau and temporoparietal lobe tau are observed years before the current threshold for Aβ-positivity. In the context of secondary prevention trials where Aβ-positivity at current thresholds is required for study inclusion, tau levels in these participants would already have been increasing for several years, likely more. The finding that temporoparietal tau starts to increase prior to other regions is in accordance with 18F-flortaucipir studies on other populations. Cross-sectional studies showed early tau deposition in cognitively healthy elderly (with or without significant Aβ pathology) in temporal and medial parietal regions, most dominant in entorhinal and parahippocampal cortex, the amygdala and inferior temporal cortex. Longitudinal studies further suggest that cognitively healthy elderly accumulate tau in the medial temporal and medial parietal lobe, while (Aβ positive) AD dementia patients increased in tau primarily in the frontal lobe (Harrison et al., 2018). The spread of tau beyond the MTL to the parietal lobe and other regions may be a critical milestone in the progression of AD. The early changes observed in the MTL in this study coincide with a recent report of the earliest tau deposition found in medial parietal regions (precuneus and isthmus cingulate) in autosomal dominant AD (Gordon et al., 2019). Considering that a 0.2 SD increase in MTL tau can potentially be detected several years before Aβ-positivity (Fig. 5), these data support the use of primary
prevention trials against Aβ where treatment is initiated years before the current threshold for Aβ-positivity, if treatment efficacy relies on early intervention, prior to the development of tau pathology.

The initial descent in cognitive performance is estimated to occur 4–6 years before becoming Aβ+ (Fig. 6). Reduced cognitive performance has repeatedly been shown to be associated with elevated levels of Aβ (Baker et al., 2017; Donohue et al., 2017; Insel et al., 2017, 2016), even within the subthreshold range (Landau et al., 2018), in cognitively unimpaired individuals. The result that CSF tau measures started to change between regional Aβ and cognition in this study is in accordance with the theory that cognitive impairment in AD is caused primarily by tau pathology. This is also in line with other recent studies which show that cognitive impairment is more strongly related to accumulation of tau than to Aβ (Ossenkoppele et al., 2019), and that both tau and Aβ appear necessary for cognitive decline (Sperling et al., 2019). The ordering of the responses coincides with the magnitude of the effect sizes at the time of Aβ-positivity (Table 1), suggesting that initial changes in the responses continue to change in parallel through to the time of Aβ-positivity, without any major differences in acceleration.
This study has several limitations. Tau PET data were available for only a subsample of the data, limiting comparisons to a small cross-section of the full ADNI data set. More data, especially longitudinal data in participants in the earliest stages of Aβ changes, will be required for more precise estimates of TFA+ as well as more precise change point estimates. Additional longitudinal Aβ information over longer periods of time will also be required to evaluate to what degree a participant may drift from their assumed quantile of accumulation. These analyses lack the power and precision to place the temporal and parietal tau regions in a particular order with confidence, but instead demonstrate that widespread tau is increasing years before Aβ-positivity. The ADNI CU, MCI, and AD cohorts are also age matched. The AD patients, on average, have dementia by age 75, while the participants in the CU cohort who may eventually develop AD, are unlikely to do so for many years, possibly decades. By design, these cohorts with age matched groups are therefore on systematically different disease trajectories with respect to age. If earlier onset is associated with a more aggressive form of the disease, then the AD cohort may have the most aggressive form while the CU cohort, the least aggressive. If the developing Aβ pathology in the ADNI CU cohort represents a less aggressive disease process compared with a more typical AD process, the estimates reported here could be conservative and biased toward later time estimates for downstream events. The ADNI MCI cohort may represent a more typical trajectory with respect to downstream events along the Aβ pathological timeline. These differences in disease trajectories are apparent from the cohort estimates in Figs. 2–5. Additionally, the change point estimates are influenced by both biological variation and measurement error, which varies from marker to marker. Change points in measures with high variability in the “normal” range and excess measurement error may require additional biological change to detect, despite an earlier, real increase in pathology. ADNI participants are highly educated on average, reducing generalizability to some degree. The associations between increasing Aβ pathology and downstream changes, including increased tau pathology reported here do not imply causality. It remains unknown whether and to what degree downstream pathological changes can be directly attributed to the accumulation of Aβ. Only studies with experimental interventions against Aβ-pathology, with clear verification of target engagement, can be used to show causal relationships between Aβ-deposition and putative downstream events. While TFA+ appears reasonably predictive, especially in proximity to the threshold for Aβ-positivity, longer follow-up is needed to validate its accuracy at very early and late-stage Aβ accumulation.

Longitudinal information is required to evaluate how quickly an individual’s pathophysiological changes are occurring and to accurately characterize their disease trajectory. Analyses limited to a cross-sectional evaluation of Aβ status are naïve to the time spent with a
significant Aβ burden. Incorporating longitudinal information facilitates the estimation of the time-course of downstream events such as the spread of tau and the onset of subtle cognitive dysfunction. As the technology to measure AD pathology becomes more cost effective and non-invasive, such as plasma measures of Aβ or tau (Janelidze et al. 2020; Mielke et al., 2018; Palmqvist et al., 2019; Schindler et al., 2019), longitudinal evaluations in the context of trial-ready cohorts may greatly improve early diagnosis and expedite the execution of clinical trials in early AD.

Data availability

All data is publicly available (http://adni.loni.usc.edu/). R code will be made available on Github.

Declaration of Competing Interest

Mr. Insel, Dr. Berron and Dr. Donohue report no competing interests. Dr. Mattsson-Carlsten has been a consultant for ADNI. Dr. Hanson has acquired research support (for the institution) from Roche, GE Healthcare, Biogen, Avid Radiopharmaceuticals and Euroimun. In the past 2 years, he has received consultancy/speaker fees (paid to the institution) from Biogen and Roche.

Credit authorship contribution statement

Philip S. Insel: Conceptualization, Methodology, Software, Formal analysis, Writing - original draft, Writing - review & editing. Michael C. Donohue: Methodology, Writing - review & editing. David Berron: Methodology, Writing - review & editing. Oskar Hanson: Writing - review & editing, Funding acquisition. Niklas Mattsson-Carlsten: Methodology, Writing - review & editing. Funding acquisition.

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References


Paper III
Determining clinically meaningful decline in preclinical Alzheimer disease

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Abstract

Objective
To determine the time required for a preclinical Alzheimer disease population to decline in a meaningful way, use estimates of decline to update previous clinical trial design assumptions, and identify factors that modify β-amyloid (Aβ)–related decline.

Methods
In 1,120 cognitively unimpaired individuals from 3 international cohorts, we estimated the relationship between Aβ status and longitudinal changes across multiple cognitive domains and assessed interactions between Aβ and baseline factors. Power analyses were performed to explore sample size as a function of treatment effect.

Results
Cognitively unimpaired Aβ+ participants approach mild cognitive impairment (MCI) levels of performance 6 years after baseline, on average. Achieving 80% power in a simulated 4-year treatment trial, assuming a 25% treatment effect, required 2,000 participants/group. Multiple factors interacted with Aβ to predict cognitive decline; however, these findings were all cohort-specific. Despite design differences across the cohorts, with large sample sizes and sufficient follow-up time, the Aβ+ groups declined consistently on cognitive composite measures.

Conclusions
A preclinical AD population declines to the cognitive performance of an early MCI population in 6 years. Slowing this rate of decline by 40%–50% delays clinically relevant impairment by 3 years—a potentially meaningful treatment effect. However, assuming a 40%–50% drug effect highlights the difficulties in preclinical AD trial design, as a more commonly assumed treatment effect of 25% results in a required sample size of 2,000/group. Designers of preclinical AD treatment trials need to prepare for larger and longer trials than are currently being considered. Interactions with Aβ status were inconsistent and not readily generalizable.
Glossary

$\beta$ = $\beta$-amyloid; AD = Alzheimer disease; ADNI = Alzheimer’s Disease Neuroimaging Initiative; AIBL = Australian Imaging, Biomarkers & Lifestyle; AIC = Akaike information criterion; BioFINDER = Biomarkers for Identifying Neurodegenerative Disorders Early and Reliably; CDR = Clinical Dementia Rating; CDR-SB = CDR sum of boxes; dADAS = Delayed Word Recall from the Alzheimer’s Disease Assessment Scale–Cognitive Subscale; dMemory = Logical Memory Delayed Recall; MCI = mild cognitive impairment; MMSE = Mini-Mental State Examination; OR = odds ratio; PACC = Preclinical Alzheimer’s Cognitive Composite; PiB = Pittsburgh compound B; SUVR = standardized uptake value ratio; Trails B = Trail-Making Test B.

To effectively alter the course of Alzheimer disease (AD), interventions may need to occur during the preclinical stage of the disease, before the onset of clinical symptoms. Demonstrating that treatments are effective during the preclinical stage will require understanding the magnitude of early $\beta$-amyloid ($A\beta$)-related cognitive decline in cognitively unimpaired adults. Defining meaningful decline will help determine the time frame for subtle cognitive changes to progress to incipient functional decline and to identify an optimal treatment window.

The association between $A\beta$ status and cognition in preclinical AD varies widely. The design of the A4 study, the first clinical trial in preclinical AD, was based on early estimates of $A\beta$-related decline using the Alzheimer’s Disease Neuroimaging Initiative (ADNI) and the Australian Imaging, Biomarkers & Lifestyle (AIBL) Study. The effect of $A\beta$ on cognitive decline in AIBL was 4-fold the magnitude of the effect in ADNI, highlighting an inconsistent picture of early cognitive decline and uncertain implications for powering a trial in early AD. Understanding how sampling variation and study design features influence estimates of cognitive decline will optimize the design of trials in preclinical AD.

The aims of this study were to harmonize several large studies in order to (1) determine the time required for a preclinical AD population to decline in a clinically meaningful way, (2) characterize how decline differs by cognitive domain, (3) update previous study design assumptions regarding sample size, power, and the required treatment effect, and (4) identify factors that modify $A\beta$-related decline.

Methods

Standard protocol approvals, registrations, and patient consents

This study was approved by the institutional review boards of all of the participating institutions. Informed written consent was obtained from all participants at each site.

Participants

Participants from each of the cohorts ADNI, AIBL, and the Swedish Biomarkers for Identifying Neurodegenerative Disorders Early and Reliably (BioFINDER) study were included if they were classified as cognitively normal at baseline, were tested for $A\beta$ biomarkers (using either CSF or PET), and were followed longitudinally with neuropsychological examinations. Participants were excluded from any of the 3 studies if they had a major neurologic or psychiatric illness or a history of substance abuse. In addition, ADNI participants were excluded if the screening MRI showed evidence of infection, infarction, or other focal lesions, including multiple lacunes or lacunes in a critical memory structure. MRI results were not part of the exclusionary criteria for AIBL or BioFINDER, but BioFINDER participants were excluded if they refused MRI or lumbar puncture. Detailed exclusionary criteria for ADNI can be found at adni.loni.usc.edu/wp-content/uploads/2008/07/adni2-procedures-manual.pdf and for BioFINDER at biofinder.se/biofinder_coaherts/cognitively-healthy-elderly/. We also included 305 participants enrolled into the early mild cognitive impairment (MCI) cohort in ADNI (defined by a subjective memory complaint and a delayed logical memory score of 9–11 for those with 16 or more years of education, 5–9 for 8–15 years of education, or 3–6 for 0–7 years of education, where possible scores range from 0 to 25) for a comparative analysis. The extensions of ADNI introduced the distinction of MCI into early and late MCI in the attempt to define an earlier point in time for disease detection. Late MCI refers to the original definition of MCI (performance for 1.5 SD below the normative mean), whereas in early MCI, impairment is defined as performance between 1.0 SD and 1.5 SD below the normative mean on a standard test. Because of recent evidence of an artificially low reversion rate from MCI to control in ADNI, we excluded 7 early MCI participants who consistently had a global Clinical Dementia Rating (CDR) score of zero after screening in a sensitivity analysis.

Data on memory complaints in the controls were available in AIBL and ADNI. In AIBL, participants with a memory complaint were identified by the response to the question, “Do you have difficulties with your memory?” In ADNI, the participant was required to have a significant memory concern as reported by the participant, study partner, or clinician and a score $>16$ on the first 12 items of the Cognitive Change Index.

$A\beta$ biomarkers

$A\beta$ status was defined by PET imaging if available (all AIBL and a majority of ADNI participants), and otherwise by CSF biomarkers (all BioFINDER and a small proportion of ADNI participants). PET imaging was done using $^{18}$F-florbetapir PET in ADNI and using $^{18}$F-florbetapir, $^{11}$C-Pittsburgh...
compound B (PiB), or 18F-flutemetamol PET in AIBL. Methods to acquire and process imaging data were described previously. CSF samples were collected at baseline by lumbar puncture. CSF methods have been described previously. In short, ADNI CSF samples were analyzed for CSF Aβ42 using the AlzBio3 assay (Fujirebio, Ghent, Belgium) on the xMAP Luminex platform. BioFINDER CSF samples were analyzed for CSF Aβ42 and Aβ40 using ELISA assays (ADx/EUROIMMUN AG, Lübeck, Germany). For ADNI participants, Aβ+ was defined as 18F-flortetapir PET standardized uptake value ratio (SUVR) > 1.1 (n = 381) or CSF Aβ42 < 192 ng/L (n = 62). For AIBL, Aβ+ was defined as 18F-flortetapir PET SUVR > 1.1 (n = 72), 11C-PiB PET SUVR > 1.5 (n = 201), or 18F-flutemetamol SUVR > 0.62 (n = 75). In BioFINDER, Aβ+ was defined as CSF Aβ42/Aβ40 < 0.1.

Cognitive testing

Participants were followed for up to 6 years for neuropsychological testing. In ADNI, tests were administered annually with an additional test at month 6 for most measures. In AIBL, tests were administered every 18 months. In BioFINDER, tests were administered every 2 years. The Preclinical Alzheimer’s Cognitive Composite (PACC) and its individual components were the primary outcomes compared in the 3 cohorts. This composite was developed specifically to be sensitive to early cognitive changes in AD and is being incorporated in clinical trials of disease-modifying treatments. Substitutions representing the same cognitive domain were made in the case where the original PACC components were not available or had limited follow-up in a cohort’s neuropsychological battery, following previous procedures. Visits where all components or substitutions were available were included. For ADNI, the modified PACC comprised the Mini-Mental State Examination (MMSE), Logical Memory Delayed Recall (dMemory), Trail-Making Test B (Trails B), and the Delayed Word Recall from the Alzheimer’s Disease Assessment Scale–Cognitive Subscale (dADASC). For AIBL, the PACC was constructed using the MMSE, dMemory, Digit Symbol Substitution Test, and the Delayed Recall from the California Verbal Learning Test (dCVLT). For BioFINDER, the PACC consisted of the MMSE, dADASC, and Trails B. To calculate the composite, z scores of the individual components were taken over all time points and then summed. This sum was then standardized to the mean and SD of the baseline score of the sum.

The PACC includes 2 measures of delayed memory recall; however, because only one delayed memory measure was available in BioFINDER, dADASC was given twice the weight in BioFINDER to reflect the contribution of delayed memory recall in the composite. Immediate recall (logical memory for ADNI and AIBL, Alzheimer’s Disease Assessment Scale–Cognitive Subscale word recall for BioFINDER) was evaluated as a measure of baseline memory ability to predict changes in the PACC. The CDR sum of boxes (CDR-SB) was also evaluated as an outcome measure.

Statistical analysis

Longitudinal measures were modeled using generalized least squares regression assuming a compound symmetric covariance structure. To capture departures from linearity in the trajectory of the neuropsychological measures, continuous time from baseline test was parameterized using restricted cubic splines. Cubic splines are functions of polynomials allowing flexibility in the estimation of trajectories over time. Time was modeled with 3 spline knots, 2 at the boundaries and 1 at median follow-up. Differences in trajectories between Aβ+ and Aβ− groups were tested using interactions between the 2 measures for time and the group factor using likelihood ratio tests and change in the Akaike information criterion (AIC), a model selection tool. A lower value of AIC indicates a better fitting model. Baseline age was also modeled using restricted cubic splines to capture its nonlinear effect on cognition. Models included the 2 spline-estimated measures for baseline age; sex; years of education, where education was categorized as 0–12 years, 13–15 years, and 16 or more years; the interaction between Aβ status and the 2 measures for time; and the main effects for Aβ status and time.

We also evaluated interactions between Aβ status and demographics (baseline age, sex, education), APOE (presence of at least one ε4 allele), memory complaint, and baseline memory, and their effect on changes in the PACC. These models included all the terms described above as well as the 3-way interaction between time, Aβ status, and the demographic term. The interaction with age was evaluated using the 2 spline-estimated measures.

To estimate power for hypothetical clinical trials, mixed models of repeated measures were used to estimate the variance components of the change from baseline in the PACC for the Aβ+ subjects in each cohort. To mirror current preclinical trial design, Aβ+ subjects with very high cognitive scores (dMemory > 15 for ADNI [n = 32] and AIBL [n = 12] and dADASC ≥ 8 in BioFINDER [n = 29]) were excluded in order to remove “supernormals.” This was done to mitigate the inclusion of participants with little or no sign of near-term decline in order to increase the likelihood of decline in the placebo group and improve power. Model estimates were then used to calculate the power for 4- and 6-year clinical trials, assuming a range of sample sizes and drug effects, a 6-month visit interval, and a 30% dropout rate. Individual cohort estimates of change from baseline and variance were then meta-analyzed to get combined estimates of change over time.

In order to provide a context for meaningful clinical decline in the cognitively normal participants, we compared the baseline PACC scores in the normal participants to the PACC scores in the ADNI early MCI participants (stratified by Aβ status). We then evaluated the mean time for the average preclinical...
AD participant to reach the mean baseline PACC score in the early MCI groups.

Baseline associations between demographics and Aβ positivity were assessed using the Wilcoxon rank-sum test for continuous variables and a χ² test for categorical variables. Reductions of AIC > 2 and p values < 0.05 were considered significant. All analyses were done in R v3.4.3 (r-project.org). GLS models were fit using the gls function from the nlme package.

Data availability
Data from the ADNI and AIBL cohorts are publicly available. Data from BioFINDER may be requested.

Results

Cohort characteristics
A total of 443 cognitively healthy controls from ADNI, 348 from AIBL, and 329 from BioFINDER were included in the study. Aβ+ groups were older, had a higher frequency of APOE ε4 positivity, and performed significantly worse on several cognitive tests at baseline, compared to Aβ− groups, in all cohorts (table 1). The proportion of APOE ε4 positivity in the Aβ+ group was similar in BioFINDER (55%) and AIBL (53%) and lower in ADNI (44%). Education and sex were not associated with Aβ positivity in AIBL or BioFINDER; however, Aβ+ ADNI participants were more likely to be female and have less education compared to Aβ− ADNI participants. The majority of ADNI participants had 16 or more years of education, whereas the majority of both AIBL and BioFINDER participants had fewer than 16 years of education. There was no association between subjective memory complaint and Aβ status in either ADNI or AIBL (subjective memory complaint data were not available in BioFINDER).

There was considerable variability in attrition rates across the 3 cohorts. At 4 years of follow-up, ADNI retained 46% of its participants; however, dropout was not associated with age, sex, education, Aβ status, or baseline memory performance (p > 0.13). At 4 years, BioFINDER retained 69% of its participants. Women were less likely to drop out (odds ratio [OR] = 0.78, p = 0.01), participants with more education were more likely to drop out (OR = 1.35, p = 0.04), and older age was associated with increased drop out (OR = 1.28 for 1 SD increase in age, p < 0.001). AIBL retained 90% of its participants, but older age was associated with increased drop out (OR = 1.26 for 1 SD increase in age, p = 0.01).

Cognitive changes
Aβ+ participants declined significantly more on the PACC and all individual components of the PACC compared to Aβ− participants, in all 3 cohorts, with the exception of Trails B in BioFINDER (p = 0.08). Estimates and longitudinal plots of cognition are shown in figure 1. Estimates of the change from baseline, confidence intervals, and the residual SD for each visit and group are shown in table 2.

At year 4, the Aβ+ groups declined by −0.45 points on the PACC (ADNI), −0.48 points (BioFINDER), and −0.53 points (at 4½ years, AIBL) (table 2). At year 4, the Aβ− group improved 0.09 points on the PACC in ADNI and declined by −0.14 points in BioFINDER and −0.02 points in AIBL.

Clinical significance
To evaluate decline and to characterize what might be considered a clinically significant change, we compared the scores of the cognitively normal participants to the baseline scores of the early MCI participants in ADNI. The mean PACC score in Aβ− and Aβ+ early MCI participants at baseline was −1.01 and −1.30, respectively (figure 2). Six years after baseline, the estimated PACC score combined across cohorts of the preclinical AD groups was midway between the Aβ− and Aβ+ early MCI performance. Similarly, the early MCI Aβ− and Aβ+ scores at baseline on the CDRSB were 1.22 and 1.38, respectively, whereas the preclinical AD groups averaged about 1.0 at 6 years.

On each of the MMSE, delayed list learning, and executive function, the cognitively normal Aβ+ groups averaged worse scores than both MCI groups by 6 years after baseline. The cognitively normal Aβ+ groups did not approach the MCI groups’ delayed logical memory scores by 6 years after baseline. Note that delayed logical memory was not available in BioFINDER.

In a sensitivity analysis, 7 early MCI participants who consistently had a global CDR of zero after screening were excluded. The reduced sample scores were slightly worse than the full MCI sample with Aβ− and Aβ+ PACC scores of −1.02 and −1.33, respectively, and CDR-SB scores of 1.23 and 1.39.

Power
Using estimates of change and variance, we calculated the power for hypothetical 4- and 6-year clinical trials for each cohort, assuming a 30% dropout rate, and various sample sizes and drug effects (figure 3). In 4-year trials, assuming a 25% drug effect, i.e., a 25% slowing of cognitive decline in the treatment group, the required sample size to reach 80% power was 2,000 per group for the estimate combining all cohorts. Assuming a larger effect size of 35%, the required sample size to reach 80% power was 1,000 per group on average.

In 6-year trials, assuming a 25% drug effect, the required sample size to reach 80% power was about 600 per group for the estimate combining all cohorts. Assuming a 35% effect size, the required sample size to reach 80% power was 300 per group on average.

Aβ interactions
The interactions between Aβ status and baseline factors to predict cognitive decline on the PACC were also assessed. Plots of the amyloid groups at different levels of the significant interacting factors, p values, and the change in AIC are shown in figure 4. In AIBL, there were significant interactions between...
### Table 1 Baseline characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Aβ+</th>
<th>Aβ−</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADNI</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Years of follow-up</td>
<td>N = 165</td>
<td>N = 278</td>
<td></td>
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<tr>
<td></td>
<td>4.1 (2.8)</td>
<td>4.3 (2.9)</td>
<td>0.28</td>
</tr>
<tr>
<td>Age</td>
<td>75.1 (5.5)</td>
<td>73.3 (5.9)</td>
<td>0.001</td>
</tr>
<tr>
<td>Female, n (%)</td>
<td>99 (60)</td>
<td>132 (47.5)</td>
<td>0.01</td>
</tr>
<tr>
<td>Education, y</td>
<td>0.77</td>
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<td></td>
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<tr>
<td>0–12</td>
<td>18 (10.9)</td>
<td>25 (9.0)</td>
<td></td>
</tr>
<tr>
<td>13–15</td>
<td>46 (27.9)</td>
<td>42 (15.1)</td>
<td></td>
</tr>
<tr>
<td>16+</td>
<td>101 (61.2)</td>
<td>211 (75.9)</td>
<td></td>
</tr>
<tr>
<td>Memory complaint, n (%)</td>
<td>42 (25.5)</td>
<td>64 (23)</td>
<td>0.64</td>
</tr>
<tr>
<td>APOE e4+, n (%)</td>
<td>73 (44.2)</td>
<td>53 (19.1)</td>
<td>&lt;0.001</td>
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<td>MMSE</td>
<td>29.1 (1.1)</td>
<td>29.0 (1.2)</td>
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<tr>
<td>dMemory</td>
<td>12.8 (3.4)</td>
<td>13.4 (3.2)</td>
<td>0.05</td>
</tr>
<tr>
<td>dADASc</td>
<td>7.0 (1.8)</td>
<td>7.2 (1.8)</td>
<td>0.33</td>
</tr>
<tr>
<td>Trails B</td>
<td>93.8 (44.4)</td>
<td>79.7 (39.4)</td>
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<td>BioFINDER</td>
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<td>N = 244</td>
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<td>Years of follow-up</td>
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<td>3.6 (1.7)</td>
<td>0.55</td>
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<tr>
<td>Age</td>
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<td>73.3 (5.0)</td>
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<td>Female, n (%)</td>
<td>56 (65.9)</td>
<td>142 (58.2)</td>
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</tr>
<tr>
<td>Education, y</td>
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<tr>
<td>0–12</td>
<td>51 (60.0)</td>
<td>146 (59.8)</td>
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<td>13–15</td>
<td>20 (23.5)</td>
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<td>40 (16.4)</td>
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<tr>
<td>Memory complaint, n (%)</td>
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<td>—</td>
<td>—</td>
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<td>APOE e4+, n (%)</td>
<td>46 (54.8)</td>
<td>46 (19)</td>
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<td>MMSE</td>
<td>29.0 (0.9)</td>
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<td>0.24</td>
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<tr>
<td>dADASc</td>
<td>7.4 (2.2)</td>
<td>8.2 (1.8)</td>
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<td>Trails B</td>
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<td>AIBL</td>
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<tr>
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<td>&lt;0.001</td>
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<tr>
<td>Age</td>
<td>73.5 (7.3)</td>
<td>69.1 (6.0)</td>
<td>&lt;0.001</td>
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<td>Female, n (%)</td>
<td>49 (49)</td>
<td>136 (54.8)</td>
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<td>Education, y</td>
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<td>0–12</td>
<td>40 (40.4)</td>
<td>104 (41.9)</td>
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<td>23 (23.2)</td>
<td>42 (16.9)</td>
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<tr>
<td>Memory complaint, n (%)</td>
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<td>132 (53.4)</td>
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<tr>
<td>APOE e4+, n (%)</td>
<td>53 (53)</td>
<td>58 (23.4)</td>
<td>&lt;0.001</td>
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### Table 1 Baseline characteristics (continued)

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<th>Characteristic</th>
<th>Aβ+</th>
<th>Aβ−</th>
<th>p Value</th>
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<td>MMSE</td>
<td>28.7 (1.2)</td>
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<td>Digit symbol</td>
<td>57.9 (12.9)</td>
<td>61.3 (13.7)</td>
<td>0.05</td>
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</tbody>
</table>

Abbreviations: Aβ = β-amyloid; ADNI = Alzheimer’s Disease Neuroimaging Initiative; AIBL = Australian Imaging, Biomarkers & Lifestyle; BioFINDER = Biomarkers for Identifying Neurodegenerative Disorders Early and Reliably; dADASc = Delayed Word Recall from the Alzheimer’s Disease Assessment Scale–Cognitive Subscale; dMemory = Delayed Recall from the California Verbal Learning Test; dMemory = Logical Memory Delayed Recall; MMSE = Mini-Mental State Examination; Trails B = Trail-Making Test B.

Aβ and education, APOE ε4 positivity, and baseline memory. The only significant interaction in ADNI was between Aβ and sex, and the only significant interaction in BioFINDER was between Aβ and age. There were no significant interactions between Aβ and subjective memory complaint (ADNI: p = 0.56, AIBL: p = 0.87, not available for BioFINDER).

### Discussion

The main findings of this study are (1) cognitively unimpaired Aβ+ participants approach early MCI cognitive performance levels on general cognition and global outcomes, delayed list recall, and executive function by 6 years after baseline; (2) to achieve 80% power in a simulated treatment trial assuming a 25% treatment effect, 2,000 participants/group for a 4-year trial and 600 participants/group for a 6-year trial are required; (3) several baseline factors interacted with Aβ status to predict decline on the PACC including APOE e4 positivity, memory, and education in AIBL; age in BioFINDER; and sex in ADNI, although these findings were all cohort-specific; (4) despite considerable design differences across the cohorts, with large sample sizes and sufficient follow-up time, the cognitively unimpaired Aβ+ groups declined consistently on cognitive composites; (5) Aβ+ groups declined significantly faster on all cognitive tests in all cohorts, with the exception of Trails B in BioFINDER, where the Aβ+ group declined marginally faster (p = 0.08), compared to the Aβ− group.

A key question for preclinical AD trials is how to define meaningful outcomes that will support use of therapeutic interventions in people who may remain asymptomatic for many years even without treatment. Traditional AD dementia trials are frequently powered to detect a several-point difference on a global cognitive score (e.g., Alzheimer’s Disease Assessment Scale–Cognitive Subscale), as well as a global/functional co-primary outcome to establish clinical meaningfulness. Post hoc analyses of the first large trials of solanezumab in patients with mild AD showed a 34% reduction of cognitive decline and a 17% reduction of functional...
However, these effects were not replicated in a subsequent randomized trial, which failed to show a significant treatment effect, with only an 11% reduction of cognitive decline and 15% reduction of functional decline. In preclinical AD, the cognitive decline observed over 3–4 years is subtle, and is typically accompanied by little or no functional decline. However, it has not been clarified what degree of decline would warrant classification as meaningful decline. To benchmark the magnitude of cognitive decline to a measure of clinical meaningfulness, we compared the scores of the cognitively unimpaired participants to those classified as early MCI—agroup with incipient functional decline. The separation between these groups was just over 1 SD on the PACC, suggesting that 1 point of additional decline in Aβ+ participants compared to Aβ− participants could be taken as an approximate benchmark for clinically meaningful decline. Combining results across cohorts shows the average Aβ+ participant to have the same PACC score at 6 years post baseline as the average patient with early MCI had at baseline (figure 2). Aβ+ participants also reached MCI level performance at 6 years on the other cognitive outcomes, with the exception of delayed logical memory. Possible explanations for this exception include that this measure was used as inclusion criterion for enrollment. This measure was also not available in BioFINDER, the cohort demonstrating the poorest scores on all measures by the end of follow-up. Finally, delayed logical memory demonstrated a clear practice effect in
the Aβ− group (figure 2), with the cognitively unimpaired participants taking this test 6 times over follow-up, compared to one time for the MCI participants.

Based on the PACC estimates, a treatment effect of 40%–50% would be required to delay the cognitive decline of a group of Aβ+ participants from reaching the 1 SD milestone by 3 years. Delaying the cognitive decline equivalent to the level of the average early MCI patient by 3 years may be a clinically meaningful treatment effect. But 40%–50% is a large treatment effect and highlights the difficulties in preclinical AD trial design. However, the observation that clinically meaningful decline is reached within 6 years offers strong support for the use of a cognitive composite in trials that are shorter than 6 years, since short term cognitive decline can be conceptualized as a proxy for downstream functional changes. With meaningful continuous cognitive changes occurring prior to an MCI diagnosis, these results, as well as recent reports,36 argue against the use of a time-to-MCI endpoint in preclinical AD trials.

The estimated sample size or trial length requirements are sobering. Previously reported sample size and drug effect requirements of 500/group with a 30%–50% effect size in a 3-year trial were optimistic and based on approximately 20% of the data available in this study.10 In order to reliably achieve 80% power for a modest, real-world effect size of 20%–30%, investors in AD research for therapeutics development will have to prepare to support larger and longer trials than are currently envisaged.

There were several significant interactions between Aβ status and baseline factors. However, no interaction was observed in more than one cohort. In AIBL, the combination of Aβ status and low education, APOE ε4 positivity, or low baseline memory all led to increased rates of decline on the PACC. Decline in the Aβ+ groups did not depend on APOE ε4 status in ADNI or BioFINDER; however, in AIBL, little decline was observed in Aβ+ participants who were not also APOE ε4+ (figure 4), as was reported previously.36,37 Evidence for additional risk of cognitive decline for individuals who are both Aβ+ and APOE ε4+ had been incorporated into the design of a phase 2b/3 trial in preclinical AD (clinicaltrials.gov/ct2/show/NCT02569398); however, this pattern was observed in only one of the 3 cohorts studied here. The additional decline observed in the Aβ+ participants who also had low baseline memory in AIBL is consistent with previous reports.38 Still, despite wide separation at baseline, high and low baseline memory (and also high and low education) groups declined in parallel over time in both ADNI and BioFINDER. The lack of replicability of these interactions across cohorts suggests that if there are true underlying effects of these baseline factors that modify the Aβ/cognition relationship, they are mild, or they depend on other/complex interactions. Another possibility is that their identification was the consequence of type I error, although the strength of the associations in AIBL (but not ADNI, reported previously39 or BioFINDER) would survive a Bonferroni correction. Our findings caution against relying on interactions between Aβ and demographic/clinical factors when selecting participants for preclinical AD trials.

There were considerable design differences among the 3 study cohorts including differences in geographic region, cognitive measures, visit frequency, and sampling characteristics.

Table 2 PACC: Change from baseline, 95% CI, and residual SD estimates

<table>
<thead>
<tr>
<th>Study</th>
<th>Month</th>
<th>N</th>
<th>Aβ+ Estimate</th>
<th>Aβ− Estimate</th>
<th>Difference (Δest)</th>
<th>95% CI</th>
<th>Residual SD (σ)</th>
<th>Δest/σ</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADNI</td>
<td>12</td>
<td>128</td>
<td>0.01</td>
<td>0.08</td>
<td>−0.08</td>
<td>−0.16 to 0.01</td>
<td>0.89</td>
<td>−0.09</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>146</td>
<td>−0.05</td>
<td>0.14</td>
<td>−0.18</td>
<td>−0.32 to −0.04</td>
<td>0.89</td>
<td>−0.21</td>
</tr>
<tr>
<td></td>
<td>36</td>
<td>62</td>
<td>−0.20</td>
<td>0.14</td>
<td>−0.34</td>
<td>−0.50 to −0.18</td>
<td>0.91</td>
<td>−0.37</td>
</tr>
<tr>
<td></td>
<td>48</td>
<td>70</td>
<td>−0.45</td>
<td>0.09</td>
<td>−0.54</td>
<td>−0.72 to −0.37</td>
<td>1.03</td>
<td>−0.53</td>
</tr>
<tr>
<td></td>
<td>60</td>
<td>31</td>
<td>−0.76</td>
<td>0.02</td>
<td>−0.78</td>
<td>−1.00 to −0.56</td>
<td>1.15</td>
<td>−0.68</td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>32</td>
<td>−1.12</td>
<td>0.08</td>
<td>−1.03</td>
<td>−1.35 to −0.72</td>
<td>1.32</td>
<td>−0.78</td>
</tr>
<tr>
<td>BioFINDER</td>
<td>24</td>
<td>75</td>
<td>−0.02</td>
<td>0.04</td>
<td>0.02</td>
<td>−0.16 to 0.20</td>
<td>0.70</td>
<td>0.03</td>
</tr>
<tr>
<td></td>
<td>48</td>
<td>55</td>
<td>−0.48</td>
<td>0.14</td>
<td>−0.34</td>
<td>−0.56 to −0.12</td>
<td>0.83</td>
<td>−0.41</td>
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<tr>
<td></td>
<td>72</td>
<td>15</td>
<td>−1.25</td>
<td>0.26</td>
<td>−0.99</td>
<td>−1.40 to −0.57</td>
<td>1.29</td>
<td>−0.77</td>
</tr>
<tr>
<td>AIBL</td>
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<td>95</td>
<td>−0.20</td>
<td>0.03</td>
<td>−0.17</td>
<td>−0.29 to −0.05</td>
<td>0.78</td>
<td>−0.22</td>
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<tr>
<td></td>
<td>36</td>
<td>81</td>
<td>−0.38</td>
<td>0.04</td>
<td>−0.34</td>
<td>−0.52 to −0.17</td>
<td>0.83</td>
<td>−0.41</td>
</tr>
<tr>
<td></td>
<td>54</td>
<td>74</td>
<td>−0.53</td>
<td>0.02</td>
<td>−0.51</td>
<td>−0.69 to −0.34</td>
<td>1.04</td>
<td>−0.49</td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>60</td>
<td>−0.66</td>
<td>0.02</td>
<td>−0.68</td>
<td>−0.88 to −0.48</td>
<td>0.98</td>
<td>−0.70</td>
</tr>
</tbody>
</table>

Abbreviations: Aβ = β-amyloid; ADNI = Alzheimer’s Disease Neuroimaging Initiative; AIBL = Australian Imaging, Biomarkers & Lifestyle; BioFINDER = Biomarkers for Identifying Neurodegenerative Disorders Early and Reliably; CI = confidence interval; PACC = Preclinical Alzheimer’s Cognitive Composite.
Despite these differences, the estimates of decline observed on the PACC in the Aβ+ groups at 4 years were remarkably similar: −0.45 points in ADNI, −0.48 in BioFINDER, and −0.53 (at 4½ years) in AIBL (table 2). Where the cohorts differed was in the change in the Aβ− group: 0.09 in ADNI, −0.14 in BioFINDER, and −0.02 in AIBL. The lower power estimate for BioFINDER for a clinical trial can be traced back to the additional decline observed in the Aβ− group, which may be due in part to including participants with presence of cerebrovascular pathology such as white matter lesions (not excluded from BioFINDER, but may have been excluded from ADNI).40,41 Cognitive reserve may also play a role, given the lower levels of education in BioFINDER compared to both ADNI and AIBL.

The Aβ group trajectories on the PACC were similar, though there was variation in the shape of the trajectories for some of the individual components. One design feature that may influence trajectory differences is test frequency. ADNI participants were tested every 6 months over the first year and every year thereafter, whereas AIBL participants were tested every 18 months and BioFINDER, every 24 months. The increased test frequency and higher levels of education in ADNI may have contributed to a tendency to improve over time as seen in dMemory (figure 1). Despite this variation in dMemory slope, Aβ− group separation over time was preserved in ADNI and AIBL. For delayed list learning, all Aβ− groups remained stable, and all Aβ+ groups showed similar decline over the total follow-up time. Combining individual components into the composite seemed to mitigate individual domain trajectory differences (figure 2). Overall, the Aβ groups across all 3 cohorts started to diverge reliably around 3 years after baseline.

One of the main limitations of this study is the variation of available measures used to construct the composite cognitive scores (i.e., the PACC) in each of the cohorts. While we included the domains represented in the original PACC, it remains unclear how these substitutions may affect the estimates of Aβ-related cognitive decline. Another limitation is
that with strict exclusionary criteria, the participants in these studies have few comorbidities, lack diversity, and do not mirror the general population. Clinical trials frequently use similar exclusionary criteria and may also lack generalizability. An additional limitation to all studies trying to inform disease-modifying AD trials is that without any information regarding potential effects of treatments, the power to detect a hypothetical effect is speculative.

Average cognitively normal Aβ+ participants approach early MCI cognitive performance levels 6 years after baseline. Comparing these 3 cohorts side by side demonstrates that large sample sizes and sufficiently long follow-up times result in consistent estimates of decline in preclinical AD. Despite substantial design and sampling differences, these results support the potential for internationally conducted clinical trials in preclinical AD. However, it is likely that designers of preclinical AD treatment trials will have to prepare for larger and longer trials than are currently considered.

**Author contributions**
P.S. Insel: drafting/revising the manuscript, study concept or design, analysis or interpretation of data, accepts responsibility for conduct of research and final approval, study supervision. M. Weiner: drafting/revising the manuscript, analysis or interpretation of data, accepts responsibility for conduct of research and final approval, study supervision. R.S. Mackin: drafting/revising the manuscript, analysis or interpretation of data, accepts responsibility for conduct of research and final approval. E. Mormino: drafting/revising the manuscript, accepts responsibility for conduct of research and final approval. Y.Y. Lim: drafting/revising the manuscript, analysis or interpretation of data, accepts responsibility for conduct of research and final approval, acquisition of data. E. Stomrud: data acquisition, accepts responsibility for conduct of research and final approval, acquisition of data, study supervision. S. Palmqvist: data acquisition, accepts responsibility for conduct of research and final approval, acquisition of data, study supervision. C.L. Masters: study concept or design, accepts responsibility for conduct of research and final approval, study supervision. P. Maruff: drafting/revising the manuscript, data acquisition, study concept or design, accepts responsibility for conduct of research and final approval, study supervision, obtaining funding. O. Hansson: drafting/revising the manuscript, data acquisition, study concept or design, accepts responsibility for conduct of research and final approval, study supervision, obtaining funding. N. Mattsson: drafting/revising the manuscript, data acquisition, study concept or design, analysis or interpretation of data, accepts responsibility for conduct of research and final approval, acquisition of data, study supervision, obtaining funding.
accepts responsibility for conduct of research and final approval, acquisition of data, study supervision, study funding.

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Figure 4 β-amyloid (Aβ) interactions

Interactions between Aβ group and age, education, sex, APOE, and baseline memory are shown for each cohort: (A) Alzheimer’s Disease Neuroimaging Initiative (ADNI), (B) Biomarkers for Identifying Neurodegenerative Disorders Early and Reliably (BioFINDER), (C) Australian Imaging, Biomarkers & Lifestyle (AIBL). Akaike information criterion (AIC) and p values are shown in each plot, testing for the significance of interactions in predicting Preclinical Alzheimer’s Cognitive Composite (PACC) change over time.
Alzheimer Immunotherapy Research & Development, LLC.; Johnson & Johnson Pharmaceutical Research & Development LLC.; Medpace, Inc.; Merck & Co., Inc.; Meso Scale Diagnostics, LLC.; NeuroRx Research; Novartis Pharmaceuticals Corporation; Pfizer Inc.; Piramal Imaging; Servier; Synarc Inc.; and Takeda Pharmaceutical Company. The Canadian Institutes of Health Research provided funds to support ADNI clinical sites in Canada. Private sector contributions were facilitated by the Foundation for the National Institutes of Health (www.fnih.org). The grantee organization is the Northern California Institute for Research and Education, and the study is coordinated by the Alzheimer’s Disease Cooperative Study at the University of California, San Diego. ADNI data are disseminated by the Laboratory for Neuro Imaging at the University of Southern California. A complete listing of ADNI investigators can be found at: https://adni.loni.usc.edu/wp-content/uploads/how_to_apply/ADNI_Acknowledgement_List.pdf. Partial financial support of AIBL was provided by the Alzheimer’s Association (US), the Alzheimer’s Drug Discovery Foundation, an Anonymous foundation, the Science and Industry Endowment Fund, the Dementia Collaborative Research Centres, the McCusker Alzheimer’s Research Foundation, the National Health and Medical Research Council (AUS), and the Yulgilbar Foundation, plus numerous commercial interactions supporting data collection. Details of the AIBL consortium can be found at www.AIBL.csiro.au and a list of the researchers of AIBL is provided at http://aibl.csiro.au/.

Disclosure
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Predicting diagnosis and cognition with $^{18}$F-AV-1451 tau PET and structural MRI in Alzheimer’s disease

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Abstract

Introduction: The relative importance of structural magnetic resonance imaging (MRI) and tau positron emission tomography (PET) to predict diagnosis and cognition in Alzheimer’s disease (AD) is unclear.

Methods: We tested 56 cognitively unimpaired controls (including 27 preclinical AD), 32 patients with prodromal AD, and 39 patients with AD dementia. Optimal classifiers were constructed using the least absolute shrinkage and selection operator with $^{18}$F-AV-1451 (tau) PET and structural MRI data (regional cortical thickness and subcortical volumes).

Results: $^{18}$F-AV-1451 in the amygdala, entorhinal cortex, parahippocampal gyrus, fusiform, and inferior parietal lobule had 93% diagnostic accuracy for AD (prodromal or dementia). The MRI classifier involved partly the same regions plus the hippocampus, with 83% accuracy, but did not improve upon the tau classifier.

Discussion: Optimized tau PET classifiers may diagnose AD with high accuracy, but both tau PET and structural brain MRI capture partly unique information relevant for the clinical deterioration in AD.

Keywords: Alzheimer; Atrophy; MRI; PET; Tau

1. Introduction

Alzheimer’s disease (AD) is characterized by β-amyloid (Aβ) and tau aggregation [1]. Aβ aggregation is assumed to lead to aggregation of tau, brain atrophy, and cognitive decline [2,3]. However, Aβ has limited toxicity and does not typically colocalize with changes in brain structure or function. In contrast, tau spreads within and beyond regions...
that show atrophy in AD and correlates with cognitive decline [4]. Tau is therefore suspected to be essential for the development of atrophy and cognitive decline in AD. Tau positron emission tomography (PET) has made it possible to study this by quantifying regional tau load in vivo [5–7]. One study on 40 patients with AD at the prodromal and dementia stages of the disease found that regional tau pathology was related to cognitive impairment and partly mediated by structural brain magnetic resonance imaging (MRI) measurements [8]. More data are needed to clarify to what degree tau aggregation and atrophy are independent processes and in which brain regions tau and atrophy are most critical for development of different symptoms. We therefore used the tau PET tracer $^{18}$F-AV-1451 together with structural brain MRI in healthy elderly controls and individuals with AD, including preclinical, prodromal, and dementia stage AD patients, to (1) identify brain regions where tau pathology and brain structure are most strongly associated with cognitive features of AD and (2) test for overlapping and complementary effects of tau and brain structure. We hypothesized that an optimized measure of tau would be superior over brain structure to identify AD and cognitive impairment. However, because atrophy and cognition may also be affected by other processes than tau pathology, including Lewy body pathology, vascular pathology, and TDP-43 pathology, we hypothesized that brain structure would provide some complementary information about cognition.

2. Methods

2.1. Participants

All participants were recruited from the Swedish BioFINDER (Biomarkers For Identifying Neurodegenerative Disorders Early and Reliably) study. Inclusion/exclusion criteria have been described previously [9,10]. We included 56 cognitively unimpaired (CU) individuals (including 27 Aβ-positive controls, called preclinical AD), 32 prodromal AD (Aβ-positive mild cognitive impairment [MCI]), and 39 AD dementia patients. All participants were assessed by physicians with expertise in dementia disorders. For CU participants, the inclusion criteria were as follows: age ≥60 years, Clinical Dementia Rating scale (CDR) 0, Mini–Mental State Examination (MMSE) 28-30, and fluency in Swedish. Exclusion criteria were as follows: refusal of lumbar puncture, presence of subjective cognitive impairment, significant neurologic or psychiatric disease, dementia or MCI, and presence of severe systemic illness. Prodromal AD cases were recruited from consecutively recruited patients with cognitive complaints at our memory clinic. The inclusion criteria were as follows: referred to the memory clinics due to cognitive complaints related to memory, executive, visuospatial, language praxis, or psychomotor function; MMSE 24–30; age 60–80 years; and fluency in Swedish. The exclusion criteria were as follows: cognitive impairment that without doubt could be explained by another condition (other than prodromal dementia), fulfillment of criteria for dementia, severe somatic disease, and refusing lumbar puncture or neuropsychological investigation. The classification of MCI (rather than subjective cognitive impairment) was based on an extensive neuropsychological battery and the assessment of a senior neuropsychologist. Patients with a clinical syndrome of AD dementia met the DSM-III-R criteria for dementia [11] and the NINCDS-ADRDA criteria for AD [12].

Informed written consent was obtained from all patients. All procedures were approved by the Regional ethics committee at Lund University, the Radiation protection committee at Skåne University Hospital, and the Swedish Medical Products Agency.

2.2. Cognitive measures

The cognitive battery used in this study included measures representing global cognition (MMSE, measured on a scale from 0 to 30, with 30 being least impaired); episodic memory (immediate and delayed wordlist recall tests from the Alzheimer’s Disease Assessment Scale–cognitive subscale [ADAS-cog], measured on scales from 0 to 10, with 0 being least impaired); processing speed/attention (Trail Making A, measured in seconds to completion of task, lower numbers indicating less impairment); and semantic memory/executive function (category [animal] fluency, measured in number of items listed, greater numbers indicating less impairment).

2.3. CSF biomarkers

Lumbar CSF sampling was done following the Alzheimer’s Association Flow Chart [13]. Samples were stored in 1-mL polypropylene tubes at −80°C until analysis. ELISA was used for analysis of CSF Aβ$_{42}$ (INNOTEST; Fujirebio, Ghent, Belgium). Aβ-positivity was defined as CSF Aβ$_{42}$ < 650 ng/L [9]. All analyses were performed by board-certified laboratory technicians who were blinded for clinical data and diagnoses.

2.4. Magnetic resonance imaging

T1-weighted MRI was performed on 3T MR scanners (Siemens Tim Trio 3T and Siemens Skyra; Siemens Medical Solutions, Erlangen, Germany), producing a high-resolution anatomical MP-RAGE image (TR = 1950 ms, TE = 3.4 ms, 1 mm isotropic voxels, and 178 slices). Cortical reconstruction and volumetric segmentation were performed with the FreeSurfer (v5.3) image analysis pipelines (http://surfer.nmr.mgh.harvard.edu). Briefly, the MP-RAGE images underwent correction for intensity homogeneity [14], removal of nonbrain tissue [15], and segmentation into gray matter and white matter with intensity gradient and connectivity among voxels [16–19]. Cortical thickness was measured as the distance from the gray matter/white matter boundary to the corresponding pial surface [17]. Reconstructed data
performed was performed using the Geometric Transfer Method [22]. Partial volume error correction because previously [20]. In brief, Hippocampus may also be susceptible to off-target binding because of its proximity to the choroid plexus [24]. However, we chose to include it because it is a recognized key region for structural brain changes and to facilitate comparisons between $^{18}$F-AV-1451 PET and MRI data.

### 2.6. Statistical analysis

The relationship between demographics and diagnosis was evaluated with Fisher’s exact test for sex and Wilcoxon-Mann-Whitney rank-sum test for age and education. Estimating the individual and joint ability of $^{18}$F-AV-1451 and MRI to predict diagnosis and cognition was done in two steps.

Step one: $^{18}$F-AV-1451 and MRI composite scores. All cognitive responses (or diagnosis) were regressed on $^{18}$F-AV-1451 retention levels or cortical thickness (or volume for hippocampus and amygdala), adjusting for demographics (age, sex, and years of education). The least absolute shrinkage and selection operator (LASSO [25]) was used for model selection and to estimate regional weights to be used to form $^{18}$F-AV-1451 and MRI composites. The LASSO selects important predictors by shrinking the individual coefficients toward zero. The coefficients of covariates that do not provide additional predictive information are shrunk to zero, resulting in parsimonious and interpretable models. The LASSO is well suited to handle large numbers of highly correlated variables such as imaging regions of interest. Ten-fold cross-validation was used to tune the amount of shrinkage. Models were subsequently fit on all data using the cross-validated penalty parameter.

Step two: Predictive value of $^{18}$F-AV-1451 and MRI composite scores. All responses were regressed on the composites developed in step one. The models were summarized with regression coefficients, standard errors, Wald test $P$ values, and the Akaike Information Criterion (AIC) [26]. The predictive ability of each imaging modality was summarized with classification accuracy for diagnosis and $R^2$ for cognitive responses. Ninety-five percent confidence intervals were estimated using jackknife estimated standard errors. Finally, all responses were regressed on both $^{18}$F-AV-1451 and MRI composites simultaneously to estimate the joint predictive ability of both modalities, as well as the adjusted regression coefficients, standard errors, and $P$ values. The reduction of the regression coefficients after adjustment was reported along with 95% confidence intervals. For prediction of diagnosis, we also present (for comparison) the accuracy of a priori selected individual regions (inferior temporal lobe for $^{18}$F-AV-1451 and hippocampal volume for MRI).

All analyses were done in R v3.3.2 (www.r-project.org).

### 3. Results

One hundred twenty-seven participants including 56 CU controls, 32 patients with prodromal AD, and 39 patients with AD dementia were examined (Table 1). All prodromal AD and AD dementia participants and 27 controls (preclinical AD) were $A_\beta^+$. For models of diagnosis, we compared all CU with the combined group of prodromal AD and AD dementia patients. The prodromal/dementia AD patients were younger on average than the CU (72.5 years vs. 74.7 years, $P = .04$) and had a higher proportion of apolipoprotein E (APOE) $e_4$ positivity (defined as the presence of one or two APOE $e_4$ alleles; 79% vs. 43%, $P < .01$). There was no difference in education (11.9 years vs. 12.2 years, $P = .76$) and a borderline significant difference in sex (63% vs. 46% male, $P = .07$).

For models of cognition, we included all preclinical AD, prodromal AD, and AD dementia participants (98 persons, with 54 males, average age 73.0 years, average education 11.9 years, 75% APOE $e_4$ positivity).

#### 3.1. $^{18}$F-AV-1451 tau PET

The $^{18}$F-AV-1451 signal was increased in prodromal AD and AD dementia in several regions throughout the temporal, parietal, frontal, and occipital lobes (see Fig. 1 for selected regions). The optimal $^{18}$F-AV-1451 classifier was 93% accurate in classifying AD (prodromal AD and AD dementia) versus CU (95% CI: 89% to 97%). The regions...
selected for classification were the amygdala, the parahippocampal gyrus, the entorhinal cortex (ERC), the fusiform cortex, and the inferior parietal lobule (Fig. 2). The a priori selected individual region inferior temporal cortical had 89% accuracy (95% CI: 80% to 98%).

Within Aβ+ participants with preclinical or clinical AD, 18F-AV-1451 was strongly associated with all cognitive responses (P < .001 for all responses). LASSO selected different regions for each cognitive test (Fig. 3). The ERC and middle temporal gyri were selected for MMSE. The parahippocampal gyrus and the ERC were selected for immediate recall. The amygdala, parahippocampal gyrus, temporal pole, ERC, and fusiform cortex were selected for delayed recall. The banks of the superior temporal sulcus, inferior temporal gyrus, lateral occipital cortex, and inferior parietal lobule were selected for Trail Making A. The inferior temporal gyrus, ERC, parahippocampal gyrus, and middle temporal gyri were selected for category fluency. The estimated regression coefficients from models predicting each cognitive response with weighted tau composites are shown in Table 2.

3.2. Magnetic resonance imaging

Structural MRI was 83% accurate in classifying participants as AD (prodromal AD and AD dementia) versus CU (95% CI 68% to 98%). The main regions selected to classify diagnosis were the ERC, hippocampus, and fusiform gyrus (Fig. 2). The a priori selected individual region hippocampus had 76% accuracy (95% CI: 60% to 92%).

Within Aβ+ participants, structural MRI was also strongly associated with all cognitive scores (P < .001 for all responses). The LASSO selected different regions for the cognitive tests (Fig. 3). The ERC, the banks of the superior temporal sulcus, and inferior parietal lobule were selected for MMSE. The parahippocampal gyrus, ERC, and the inferior parietal lobule were selected for immediate recall. The hippocampus, ERC, amygdala, parahippocampal gyrus, banks of the superior temporal sulcus, and the inferior parietal lobule were selected for delayed recall. Several regions were selected for Trail Making A, with the inferior temporal gyrus, fusiform gyrus, and isthmus cingulate being the most influential regions. The ERC was selected for category fluency. The estimated regression coefficients from models predicting each cognitive response with weighted brain MRI composites are shown in Table 2.

3.3. Competing and complementary predictive information: MRI and 18F-AV-1451

The effect estimates from regression models predicting diagnosis or cognition with (1) 18F-AV-1451 only, (2) MRI only, and (3) 18F-AV-1451 and MRI are shown in Table 2. 18F-AV-1451 showed strongest associations with delayed recall (R² = 0.48), followed by immediate recall (R² = 0.41), MMSE (R² = 0.36), category fluency (R² = 0.33), and Trail Making A (R² = 0.23). MRI had similar strength of associations with delayed recall (R² = 0.48), MMSE (R² = 0.35), immediate recall (R² = 0.34), category fluency (R² = 0.29), and Trail Making A (R² = 0.22). Results from the three models show the change in the effect estimate when adjusting for the other imaging modality. The percent reduction of the regression estimate and a 95% confidence interval are shown in Table 2, as well as AIC values for each model. The estimates for 18F-AV-1451 were reduced between −2% (for diagnosis) and 43% (for MMSE) when adjusting for MRI. Reduction of MRI estimates ranged from 35% (for diagnosis) to 49% (for immediate recall) when adjusting for tau. AIC selected the models (ΔAIC when comparing two models > 2 favors the model with smallest AIC) with both 18F-AV-1451 and MRI to predict diagnosis and all cognitive responses (Table 2). Adjusted and unadjusted regression estimates and confidence intervals are shown in Fig. 4.

Table 1

Demographics

<table>
<thead>
<tr>
<th></th>
<th>Aβ− CU</th>
<th>Aβ+ CU (preclinical AD)</th>
<th>Aβ+ MCI (prodromal AD)</th>
<th>Aβ+ AD dementia</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>29</td>
<td>27</td>
<td>32</td>
<td>39</td>
<td></td>
</tr>
<tr>
<td>Age, y</td>
<td>75.0 (5.5)</td>
<td>74.4 (7.6)</td>
<td>72.5 (6.9)</td>
<td>76.2 (6.8)</td>
<td>.25</td>
</tr>
<tr>
<td>Male, N (%)</td>
<td>17 (59%)</td>
<td>9 (33%)</td>
<td>21 (66%)</td>
<td>24 (62%)</td>
<td>.063</td>
</tr>
<tr>
<td>Education, y</td>
<td>12.7 (3.6)</td>
<td>11.7 (4.2)</td>
<td>11.9 (3.1)</td>
<td>11.9 (3.8)</td>
<td>.66</td>
</tr>
<tr>
<td>APOE e4, −/+ (%)</td>
<td>227/24%</td>
<td>10/17 (63%)</td>
<td>4/28 (88%)</td>
<td>10/26 (72%)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>MMSE</td>
<td>29.0 (1.1)</td>
<td>29.3 (1.0)</td>
<td>26.5 (2.5)</td>
<td>21.7 (4.6)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Immediate Recall</td>
<td>2.6 (1.2)</td>
<td>2.3 (1.1)</td>
<td>4.9 (1.4)</td>
<td>6.3 (1.5)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Delayed Recall</td>
<td>1.9 (1.4)</td>
<td>2.0 (1.5)</td>
<td>5.8 (2.6)</td>
<td>8.4 (2.0)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Trail Making A</td>
<td>3.72 (0.31)</td>
<td>3.85 (0.28)</td>
<td>3.98 (0.36)</td>
<td>4.28 (0.44)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Category Fluency</td>
<td>24.7 (5.6)</td>
<td>21.6 (5.5)</td>
<td>16.3 (5.1)</td>
<td>11.1 (5.4)</td>
<td>&lt;.001</td>
</tr>
</tbody>
</table>

Abbreviations: Aβ, β-amyloid; AD, Alzheimer’s disease; APOE, apolipoprotein E; CU, cognitively unimpaired; MMSE, Mini–Mental Status Examination.

NOTE. Continuous data are mean (standard deviation). Trail Making A data are shown after natural log transformation. The combined CU (including both Aβ− and Aβ+) had mean age 74.7 (6.57) years, mean education 12.2 (3.9) years, mean MMSE 29.1 (1.1), mean immediate recall 2.4 (1.1), mean delayed recall 1.9 (1.5), mean Trail Making A 3.79 (0.30), and mean category fluency 23.2 (5.8). P values by Kruskal-Wallis test for continuous data and Fisher’s exact test for categorical data.
4. Discussion

We tested the predictive value of $^{18}$F-AV-1451 tau PET and structural MRI to identify AD and associations with cognition. The main findings were that (1) an optimized classifier that used regional $^{18}$F-AV-1451 had superior diagnostic accuracy for AD compared to brain MRI; (2) $^{18}$F-AV-1451 and MRI had overall similar strengths of associations with cognition; and (3) both $^{18}$F-AV-1451 and MRI contributed complementary information about cognitive impairment through the continuum from preclinical to prodromal and dementia stages of AD, with regional differences between the modalities. Several previous studies have found that biomarkers of brain structure [27,28] and tau [5,6] are associated with clinical features of AD. However, studies directly comparing these processes and their regional effects on diagnosis and cognition in AD are rare [8]. Our novel findings suggest that although tau is the most critical of the two, both $^{18}$F-AV-1451 tau PET and structural brain MRI capture partly unique information that is relevant for the clinical deterioration in AD.

With 93% accuracy, our $^{18}$F-AV-1451 classifier was excellent for identification of clinical AD. The selected regions were mainly temporal lobe regions, where tau pathology presumably occurs in early stages of AD (ERC, amygdala, fusiform, and the parahippocampal gyrus), but
also the inferior parietal lobule, which presumably is involved in later stages of the disease [7]. Note that these results are for a combination of patients with AD at the prodromal and dementia stages, compared with CU (which included 48% preclinical AD). This suggests that $^{18}$F-AV-1451 tau PET may be used as a powerful instrument for diagnosis of AD at the clinical stage, without considering the somewhat arbitrary distinction between prodromal and dementia stages.

The optimal MRI-based classifier achieved lower accuracy (83%) and partly included similar regions as the $^{18}$F-AV-1451 classifier (ERC, fusiform) plus the hippocampus, the banks of the superior temporal sulcus, and the inferior parietal lobule. These regions are similar to but not identical with a previously proposed “temporal meta-ROI” to capture AD-related atrophy (the ERC, inferior temporal, middle temporal, and fusiform [29]). The MRI classifier also included several regions with negative coefficients (meaning that greater thickness was associated with AD diagnosis), including pars opercularis, superior temporal, paracentral, and lateral orbitofrontal cortex. One interpretation of these negative coefficients is that they control for premorbid differences in cortical thickness, but it is also possible that the negative coefficients represent variability particular to this data set. The lower accuracy of MRI compared with $^{18}$F-AV-1451 may be due to the lower specificity of brain

![Fig. 2. $^{18}$F-AV-1451 and brain MRI for diagnosis of AD. The top part shows model-estimated regional weights and boxplots of composites for both $^{18}$F-AV-1451 (top row) and MRI (middle row) for the classification of diagnosis (including prodromal AD and AD dementia). The bottom part shows visualization of the model-selected regions, color-coded according to the respective model-estimated weights. Abbreviations: Aβ, β-amyloid; AD, Alzheimer’s disease; ERC, entorhinal cortex; MRI, magnetic resonance imaging; STS, superior temporal sulcus; SUVR, standardized uptake value ratio.](image-url)
Fig. 3. $^{18}$F-AV-1451, brain MRI and cognition. The top part shows model-estimated regional weights in barplots for $^{18}$F-AV-1451 (left) and brain MRI (right) to predict different cognitive tests in AD (including preclinical AD, prodromal AD, and AD dementia patients). For each cognitive test and modality, cognitive scores are plotted against the composites resulting from the model-estimated weights. Regression curves and 95% confidence intervals are shown in red. The bottom part shows visualization of the model-selected regions, color-coded according to the respective model-estimated weights. Abbreviations: Aβ, β-amyloid; AD, Alzheimer’s disease; ERC, entorhinal cortex; MRI, magnetic resonance imaging; STS, superior temporal sulcus; SUVR, standardized uptake value ratio.
structure because a variability in the brain structure is seen both in normal aging and due to non-AD diseases, and due to greater sensitivity of \(^{18}\)F-AV-1451 to detect subtle AD pathology.

In addition to the higher diagnostic accuracy for \(^{18}\)F-AV-1451, we found that when combining tau and brain structure, the association between \(^{18}\)F-AV-1451 and AD diagnosis was minimally affected by adjusting for MRI (increased by 2%), but the association between MRI and AD was markedly reduced by adjusting for tau (reduced by 35%). The classification accuracy for the model including both \(^{18}\)F-AV-1451 and MRI did not improve over \(^{18}\)F-AV-1451 alone, indicating that brain structure contributes little beyond tau to the identification of patients with AD.

Hippocampus was selected for the MRI classifier but not for the \(^{18}\)F-AV-1451 classifier. There are several possible explanations for this. First, the hippocampus may be sensitive to off-target binding for \(^{18}\)F-AV-1451 because of its proximity to the choroid plexus, and noise from off-target binding may reduce the importance of the hippocampus for tau quantification [24] (although we used partial volume error corrected data, which reduces the influence of off-target binding, rendering estimates less sensitive to this confounding factor). Second, tau accumulation in the hippocampal structures besides CA1 may occur quite late in AD [7], and hippocampal sparing types of AD are not rare [30]. Together, these factors likely reduce the importance of hippocampal \(^{18}\)F-AV-1451 for AD diagnosis. In contrast, hippocampal atrophy

<table>
<thead>
<tr>
<th>Response</th>
<th>(^{18})F-AV-1451</th>
<th>MRI only</th>
<th>(^{18})F-AV-1451 and MRI</th>
<th>(^{18})F-AV-1451</th>
<th>MRI only</th>
<th>(^{18})F-AV-1451 and MRI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diagnosis</td>
<td>8.48 4.84</td>
<td>2.14 2.00</td>
<td>3.66 7.66</td>
<td>4.04 2.00</td>
<td>3.66 7.66</td>
<td>4.04 2.00</td>
</tr>
<tr>
<td>MMSE</td>
<td>2.66 1.52</td>
<td>0.37 0.20</td>
<td>534.6 43.6</td>
<td>2.80 1.77</td>
<td>0.37 0.20</td>
<td>531.8 43.6</td>
</tr>
<tr>
<td>Immediate Recall</td>
<td>1.52 1.41</td>
<td>0.35 0.20</td>
<td>38.3 20.1</td>
<td>1.52 1.41</td>
<td>0.35 0.20</td>
<td>38.3 20.1</td>
</tr>
<tr>
<td>Delayed Recall</td>
<td>1.52 1.41</td>
<td>0.35 0.20</td>
<td>38.3 20.1</td>
<td>1.52 1.41</td>
<td>0.35 0.20</td>
<td>38.3 20.1</td>
</tr>
<tr>
<td>Trail Making A</td>
<td>0.20 0.20</td>
<td>0.14 0.05</td>
<td>376.9 17.0</td>
<td>0.20 0.20</td>
<td>0.14 0.05</td>
<td>376.9 17.0</td>
</tr>
<tr>
<td>Category Fluency</td>
<td>3.88 3.88</td>
<td>0.58 0.20</td>
<td>564.7 27.0</td>
<td>3.88 3.88</td>
<td>0.58 0.20</td>
<td>564.7 27.0</td>
</tr>
</tbody>
</table>

Abbreviations: A\(_b\), t-amyloid; AD, Alzheimer’s disease; AIC, Akaike Information Criterion; MRI, cognitively unimpaired; LASSO, least absolute shrinkage and selection operator; MMSE, Mini–Mental Status Examination; MRI, magnetic resonance imaging.

Fig. 4. Adjusted and unadjusted models for diagnosis and cognition. Regression estimates with and without adjustment for the other imaging modality are plotted with 95% confidence intervals (CIs) for each response (corresponding to Table 2). Signs for MRI estimates were inverted to facilitate comparisons with \(^{18}\)F-AV-1451 estimates. All models were adjusted for age, sex and education. Abbreviations: MMSE, Mini–Mental Status Examination; MRI, magnetic resonance imaging.
is a key aspect of structural brain changes in AD, except for in rare variants [31].

The a priori selected regions for \(^{18}\text{F-AV-1451}\) (inferior temporal lobe) and MRI (hippocampus) had lower accuracy than the optimal classifiers (89% vs. 93% for \(^{18}\text{F-AV-1451}\) and 76% vs. 83% for MRI), but the differences were not statistically significant. The small increase for \(^{18}\text{F-AV-1451}\) (4%) and the small/moderate increase for MRI (7%) may be clinically meaningful, although the power to detect those increases with the current sample was low.

Increased \(^{18}\text{F-AV-1451}\) retention was closely related to worse cognition in AD. The strongest association was seen for delayed recall. MRI had similar associations, but when combining \(^{18}\text{F-AV-1451}\) and MRI, \(^{18}\text{F-AV-1451}\) was sometimes more strongly related to cognition, with greater reductions of the MRI coefficient. In particular, the effect of \(^{18}\text{F-AV-1451}\) on immediate recall was reduced by 30% when adjusting for MRI, whereas the effect of MRI was reduced by 49% when adjusting for \(^{18}\text{F-AV-1451}\). However, both \(^{18}\text{F-AV-1451}\) and MRI provided partly complementary information that was not completely accounted for by the other modality, especially for general cognition (MMSE) and delayed recall. Note the difference between these associations with cognition and the associations with diagnosis described previously. When dichotomizing into diagnoses, MRI did not appear to contribute additional predictive information beyond \(^{18}\text{F-AV-1451}\). When evaluating a continuum of cognitive scores in multiple domains throughout the course from preclinical, prodromal, and dementia stages of AD, both \(^{18}\text{F-AV-1451}\) and MRI provided predictive information.

The partly independent effects of \(^{18}\text{F-AV-1451}\) on cognition may suggest that tau had effects on neuronal function and integrity, with relevance for cognition before overt atrophy was seen by MRI. It has been demonstrated before that cognitive changes may occur in parallel with or even precede atrophy measures during the course of AD [32], and according to animal studies, tau may have very early effects on neuronal activity [33]. In contrast, the relationship between MRI and cognition that was partly independent of \(^{18}\text{F-AV-1451}\) may reflect atrophy that has accelerated downstream of tau or atrophy caused by vascular disease, TDP-43 pathology, Lewy bodies, or other processes that are independent of tau [34].

The regions that were selected for cognition differed slightly from those that were selected for diagnosis and differed between \(^{18}\text{F-AV-1451}\) and MRI. For immediate recall, the optimal tau composite was sampled from temporal regions (parahippocampal, temporal pole, amygdala, ERC, fusiform), and the optimal MRI composite included partly overlapping regions (parahippocampal, amygdala, ERC, hippocampus, inferior parietal lobule, and banks of the superior temporal sulcus). This complex regional relationship between tau, atrophy, and cognition may have implications both for future research and clinical trials. For example, if tau PET and MRI were used as trial endpoints, it may be beneficial to measure longitudinal effects in modality-dependent regions to optimize chances of detecting clinically relevant effects.

One limitation is that we only used cross-sectional biomarker data. Future use of longitudinal data may give further insights into the relationships between tau, atrophy, and cognitive decline [35,36]. Another limitation is that the relative weights in the LASSO may differ depending on slight variations in the correlations between the predictors and the response, and we therefore caution against overinterpreting the ordering of the weights. Furthermore, models with a large number of predictors may be prone to overfitting. Here, we used 10-fold cross-validation to tune the LASSO penalty parameter. By splitting the data into training and test sets, the value of the penalty parameter was selected based on subjects outside the training set. The tuning of this parameter was done to maximize parsimony and prevent overfitting.

We conclude that \(^{18}\text{F-AV-1451}\) tau PET was strongly associated with AD diagnosis, with stronger associations than structural MRI. However, both \(^{18}\text{F-AV-1451}\) and structural MRI were independently associated with cognitive impairment, in a cross-sectional analysis that included the entire disease continuum from preclinical to prodromal and dementia stages of AD.

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Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jalz.2018.12.001.
1. Systematic review: We reviewed the literature using PubMed and complemented with key papers from reference lists. There are a few papers on combined brain magnetic resonance imaging and tau positron emission tomography (PET) in AD, which suggest that tau PET is more closely linked to cognition. But the results have not been conclusive, and to our knowledge, optimized classifiers for tau accumulation and brain atrophy to predict diagnosis and cognitive impairment in AD have not been tested.

2. Interpretation: In this cohort-study that included 127 participants, an optimal regional tau PET classifier had 93% diagnostic accuracy for the prodromal or dementia stages of AD, whereas an optimal brain magnetic resonance imaging classifier had 83% accuracy and did not improve upon the tau classifier. Both tau and magnetic resonance imaging were strongly (and party independently) associated with cognitive impairment through the preclinical, prodromal, and dementia stages of AD. This supports that tau PET measures provide superior information about diagnosis and brain changes relevant for cognitive decline in AD.

3. Future directions: Studies incorporating longitudinal brain structure, tau PET, and cognition are necessary to understand the spatiotemporal dynamics of these changes in AD.

References


IMPORTANCE Although the most common recent approach in Alzheimer disease drug discovery is to directly target the β-amyloid (Aβ) pathway, the high prevalence of apolipoprotein E ε4 (APOE ε4) in Alzheimer disease and the ease of identifying ε4 carriers make the APOE genotype and its corresponding protein (apoE) an appealing therapeutic target to slow Aβ accumulation.

OBJECTIVE To determine whether the ε2 allele is protective against Aβ accumulation in the presence of the ε4 allele and evaluate how age and the APOE genotype are associated with emerging Aβ accumulation and cognitive dysfunction.

DESIGN, SETTING, AND PARTICIPANTS This cross-sectional study used screening data from the Anti-Amyloid Treatment in Asymptomatic Alzheimer Disease Study (A4 Study) collected from April 2014 to December 2017 and analyzed from November 2019 to July 2020. Of the 6943 participants who were a part of the multicenter clinical trial screening visit, 4432 were adults without cognitive impairment aged 65 to 85 years who completed a fluorine 18-labeled (18F)-florbetapir positron emission tomography scan, had APOE genotype information, and had a Clinical Dementia Rating of 0. Participants who were taking a prescription Alzheimer medication or had a current serious or unstable illness that could interfere with the study were excluded.

MAIN OUTCOMES AND MEASURES Aβ pathology, measured by 18F-florbetapir positron emission tomography and cognition, measured by the Preclinical Alzheimer Cognitive Composite.

RESULTS A total of 4432 participants were included (mean [SD] age, 71.3 [4.7] years; 2634 women [59.4%]), with a mean (SD) of 16.6 (2.8) years of education and 1512 (34.1%) with a positive Aβ level. APOE ε2 was associated with a reduction in both the overall (standardized uptake value ratio [SUVR], ε24, 1.11 [95% CI, 1.08-1.14]; ε34, 1.18 [95% CI, 1.17-1.19]) and the age-dependent level of Aβ in the presence of ε4, with Aβ levels in the APOE ε24 group (n = 115; ε24, 0.005 SUVR increase per year of age) increasing at less than half the rate with respect to increasing age compared with the APOE ε34 group (n = 1295; 0.012 SUVR increase per year of age; P = .04). The association between Aβ and decreasing Preclinical Alzheimer Cognitive Composite scores did not differ by APOE genotype, and the reduced performance on the Preclinical Alzheimer Cognitive Composite in APOE ε4 carriers compared with noncarriers was completely mediated by Aβ (unadjusted difference in composite scores between ε4 carriers and noncarriers = −0.084, P = .005; after adjusting for 18F-florbetapir = −0.006, P = .85; after adjusting for 18F-florbetapir and cardiovascular scores = −0.009, P = .78).

CONCLUSIONS AND RELEVANCE These findings suggest that the protective outcome of carrying an ε2 allele in the presence of an ε4 allele against Aβ accumulation is important for potential treatments that attempt to biochemically mimic the function of the ε2 allele in order to facilitate Aβ clearance in ε4 carriers. Such a treatment strategy is appealing, as ε4 carriers make up approximately two-thirds of patients with Alzheimer disease dementia. This strategy could represent an early treatment option, as many ε4 carriers begin to accumulate Aβ in early middle age.
Age and the apolipoprotein E (APOE) genotype are among the strongest risk factors for amyloid-β (Aβ) accumulation. Rates of early Aβ accumulation are highest in APOE ε4 allele carriers and lowest in ε2 allele carriers compared with ε3-only carriers. Increasing evidence suggests that the APOE genotype and its corresponding protein (apoE) affect the pathogenesis of Alzheimer disease (AD) through multiple biological pathways, including the differential regulation of Aβ aggregation and clearance. Two-thirds of patients with AD dementia are APOE ε4 allele carriers. Although the most common recent approach in AD drug discovery is to directly target the Aβ pathway, the high prevalence of APOE ε4 in AD and the ease of identifying ε4 carriers at any age make APOE pathways an appealing therapeutic target to slow Aβ accumulation.

Two single-nucleotide variations in APOE and 3 apoE isoforms (apoE2, apoE3, and apoE4) are thought to have a substantial effect on the structure and function of apoE, including Aβ binding. From a treatment standpoint, it is unclear whether strategies that increase the “good” forms of apoE (apoE2, apoE3) or decrease the “bad” form (apoE4) would be most successful. By mimicking the biochemical properties associated with the apoE2 isoform, it may be possible to increase the Aβ clearance that is reduced with apoE4. However, a central question is whether apoE2 remains protective in the presence of apoE4. This question has been difficult to answer, in large part because the simultaneous carriage of both the ε2 and ε4 alleles is rare—approximately 2% of the population has the ε24 genotype. Even in previous meta-analyses, small sample sizes of the ε24 group precluded precise estimates of the effect of ε2 in the presence of ε4 on Aβ pathology. Consequently, questions about the potential for therapies targeting apoE, such as synthetic peptides, to reduce Aβ pathology in the presence of apoE4 remain unanswered.

Because genetic risk factors, such as carriage of APOE ε4, can be determined at birth, targeting apoE4 is of particular interest as an early treatment option. Predicting when individuals may become at increased risk for abnormal rates of Aβ accumulation will help to inform the design of primary prevention. In 4432 participants without cognitive impairment who were screened for participation in the Anti-Amyloid Treatment in Asymptomatic Alzheimer Disease (A4 Study) trial, we evaluated how the principal risk factors for AD (age and APOE genotype) were associated with early buildup of Aβ, measured by fluroine 18-labeled (18F)-florbetapir positron emission tomography (PET).

### Methods

In this cross-sectional study, data were collected from April 2014 to December 2017 and analyzed from November 2019 to July 2020. Participants screened for inclusion in the A4 Study were included in this study if they completed an 18F-florbetapir PET scan, had APOE genotype information, completed a battery of neuropsychological testing, scored between 25 and 30 on the Mini-Mental State Examination (MMSE), had a Clinical Dementia Rating of 0, and were aged between 65 and 85 years. Exclusion criteria for the A4 study have been described previously. Briefly, participants were excluded from the A4 Study if they were taking a prescription Alzheimer medication or had a current serious or unstable illness that could interfere with the study. Note that participants without evidence of brain Aβ at screening were not randomized to treatment in the A4 Study but were included in the current study regardless of their PET scan result. This study was approved by the institutional review boards of all participating institutions, and written informed consent was obtained from all participants. This study followed the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) reporting guideline.

### 18F-Florbetapir PET Imaging

β-Amyloid PET imaging in the A4 Study was done using 18F-florbetapir data, which was acquired 50 to 70 minutes postinjection. Images were realigned and averaged and then spatially aligned to a standard space template. 18F-florbetapir, sampled in a global neocortical region for Aβ, was expressed as a standardized uptake value ratio (SUVR) with a cerebellar reference region. β-Amyloid positivity was defined as participants with an 18F-florbetapir PET SUVR greater than or equal to 1.10.11,12

### Cognitive Testing

The A4 Study participants completed a neuropsychological test battery including the Preclinical Alzheimer Cognitive Composite (PACC), comprising the MMSE, the Logical Memory Delayed Recall, the Free and Cued Selective Reminding Test, and the Digit Symbol Substitution Test. To calculate the PACC, individual components were z-transformed and summed. The resulting sum was then centered on the mean and SD of the Aβ-negative group.

### Cardiovascular Risk Factors

Cardiovascular risk scores were calculated based on body mass index, systolic blood pressure, smoking status, and information gathered during an initial health assessment and physical and neurologic examination. During the initial assessment, participants were asked about the chronicity and severity of underlying health conditions. Chronicity (1 = single occurrence; 2 = intermittent; and 3 = persistent) and severity...
scores were used to calculate the cardiovascular risk score, described further in the Statistical Analysis section. During the physical and neurologic examination, participants were classified as normal or abnormal with regard to cardiac health. Cardiac values (0 = normal; 1 = abnormal) were incorporated into the cardiovascular risk score.

### Statistical Analysis

18F-Florbetapir SUVR values (both continuous and dichotomized, separately) were regressed on APOE genotype (all 6 genotypes), adjusting for age and sex. The interaction between APOE genotype and age was also assessed. Models with continuous outcomes (18F-florbetapir PET SUVRs and cognitive scores) were modeled using ordinary least-squares regression. Monotone cubic splines were used to evaluate potential nonlinearity in the associations among 18F-florbetapir PET SUVR, age, and cognition. Statistical significance of the associations between the outcome and predictors was tested using likelihood ratio tests and the Akaike information criterion (AIC). A lower value of the AIC indicates a better-fitting model. Multiple-comparison P value adjustment of the PACC components was done using a Holm correction. All models predicting 18F-florbetapir PET included age and sex. All models predicting cognition included age, sex, and years of education.

Cardiovascular risk scores were also evaluated for their association with the outcomes and the association between APOE genotype and the outcomes. Cardiovascular risk scores were calculated as the sum of z-transformed body mass index, z-transformed natural log systolic blood pressure, z-transformed product of chronicity and severity of cardiovascular symptoms from the initial health assessment, smoking status (0 = nonsmoker; 1 = smoker), and cardiac symptoms from the physical and neurologic examination (0 = normal; 1 = abnormal). The resulting sum was then z-transformed, providing a summary of cardiovascular risk with higher scores indicating more risk.

Associations between demographics and APOE genotype were assessed using a Kruskal-Wallis test for continuous variables and a χ² test for categorical variables. Changes in the AIC (ΔAIC) less than -2 and 2-sided P < .05 were considered significant. The PACC P values were obtained using the Holm-Bonferroni method. All analyses were done in R software, version 3.6.0 (R Foundation).

### Results

Of the 6943 participants who were part of the multicenter clinical trial screening visit, 4432 adults without cognitive impairment were included (2634 women [59.4%] and 1798 men [40.6%]; mean [SD] age, 71.3 [4.7] years). Individuals had mean (SD) of 16.6 (2.8) years of education, and 1512 had a positive Aβ level (34.1%). Cohort characteristics by APOE genotype are summarized in Table 1.

### Table 1. Characteristics of the Study Cohort by APOE Genotype

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>APOE ε22 (n = 25)</th>
<th>APOE ε23 (n = 449)</th>
<th>APOE ε33 (n = 2409)</th>
<th>APOE ε24 (n = 115)</th>
<th>APOE ε34 (n = 1295)</th>
<th>APOE ε44 (n = 139)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean (SD), y</td>
<td>71.1 (4.1)</td>
<td>71.9 (4.9)</td>
<td>71.3 (4.8)</td>
<td>71.5 (5.0)</td>
<td>70.7 (4.3)</td>
<td>69.8 (3.8)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Women, No. (%)</td>
<td>16 (64.0)</td>
<td>245 (54.6)</td>
<td>1455 (60.4)</td>
<td>66 (57.4)</td>
<td>771 (59.5)</td>
<td>81 (58.3)</td>
<td>.32</td>
</tr>
<tr>
<td>Race No. (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>25 (100)</td>
<td>395 (88.0)</td>
<td>2188 (90.8)</td>
<td>103 (89.6)</td>
<td>1203 (92.9)</td>
<td>128 (92.1)</td>
<td></td>
</tr>
<tr>
<td>Black</td>
<td>0</td>
<td>31 (6.9)</td>
<td>71 (2.9)</td>
<td>9 (7.8)</td>
<td>41 (3.2)</td>
<td>7 (5.0)</td>
<td></td>
</tr>
<tr>
<td>Asian</td>
<td>0</td>
<td>15 (3.3)</td>
<td>117 (4.9)</td>
<td>2 (1.7)</td>
<td>33 (2.6)</td>
<td>2 (1.4)</td>
<td></td>
</tr>
<tr>
<td>American Indian</td>
<td>0</td>
<td>1 (0.2)</td>
<td>3 (0.1)</td>
<td>0</td>
<td>4 (0.3)</td>
<td>1 (0.7)</td>
<td>.003</td>
</tr>
<tr>
<td>Native Hawaiian</td>
<td>0</td>
<td>0</td>
<td>1 (0.04)</td>
<td>0</td>
<td>1 (0.08)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Multiple</td>
<td>0</td>
<td>4 (0.9)</td>
<td>14 (0.6)</td>
<td>0</td>
<td>9 (0.7)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>0</td>
<td>3 (0.7)</td>
<td>15 (0.6)</td>
<td>1 (0.9)</td>
<td>4 (0.3)</td>
<td>1 (0.7)</td>
<td></td>
</tr>
<tr>
<td>Ethnicity, No. (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hispanic or Latino</td>
<td>0</td>
<td>8 (1.8)</td>
<td>89 (3.7)</td>
<td>6 (5.2)</td>
<td>29 (2.2)</td>
<td>6 (4.3)</td>
<td></td>
</tr>
<tr>
<td>Non-Hispanic or non-Latino</td>
<td>25 (100)</td>
<td>438 (97.6)</td>
<td>2300 (95.5)</td>
<td>108 (93.9)</td>
<td>1255 (96.9)</td>
<td>133 (95.7)</td>
<td>.21</td>
</tr>
<tr>
<td>Unknown</td>
<td>0</td>
<td>3 (0.7)</td>
<td>20 (0.8)</td>
<td>1 (0.9)</td>
<td>11 (0.8)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Education, mean (SD), y</td>
<td>17.2 (3.5)</td>
<td>16.5 (3.1)</td>
<td>16.3 (2.8)</td>
<td>16.6 (3.2)</td>
<td>16.7 (2.8)</td>
<td>16.5 (2.7)</td>
<td>.70</td>
</tr>
<tr>
<td>Cardiovascular Risk Score, mean (SD)</td>
<td>-0.06 (1.01)</td>
<td>-0.09 (0.98)</td>
<td>0.00 (0.99)</td>
<td>0.16 (0.86)</td>
<td>-0.03 (1.01)</td>
<td>-0.14 (0.96)</td>
<td>.08</td>
</tr>
<tr>
<td>Aβ positive, No. (%)</td>
<td>4 (16.0)</td>
<td>82 (18.3)</td>
<td>562 (23.3)</td>
<td>47 (40.9)</td>
<td>702 (54.2)</td>
<td>115 (82.7)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Aβ SUVR, estimated mean (95% CI)</td>
<td>1.02 (0.95-1.09)</td>
<td>1.02 (1.01-1.04)</td>
<td>1.05 (1.04-1.06)</td>
<td>1.11 (1.08-1.14)</td>
<td>1.18 (1.17-1.19)</td>
<td>1.31 (1.28-1.34)</td>
<td>&lt;.001</td>
</tr>
</tbody>
</table>

Abbreviations: Aβ, β-amyloid; APOE, apolipoprotein E; SUVR, standardized uptake value ratio.
**Figure 1. Apolipoprotein E (APOE), Age, and Fluorine 18-Labeled (18F)-Florbetapir Standardized Uptake Value Ratio (SUVR)**

**A** 18F-florbetapir PET by genotype

Boxplots of continuous 18F-florbetapir SUVRs are shown for each APOE group (A). The horizontal lines inside the boxes indicate the median; the upper and lower bounds of the boxes indicate the third and first quartiles, respectively; whiskers indicate the most extreme point no more than 1.5 times the interquartile range; and circles indicate outliers. To the right of each boxplot is the whiskers indicating the most extreme point no more than 1.5 times the interquartile range; and circles indicate outliers. To the right of each boxplot is the mean and 95% CI. The 18F-florbetapir SUVR is plotted by age for each APOE group, separately (ε22 and ε23 participants were combined because of the small sample size over age for the ε22 group) (B). The estimated probability of Aβ positivity is plotted by age for each APOE group, separately (C). Aβ indicates β-amyloid; PET, positron emission tomography. The shaded areas in B and C indicate 95% CIs.

**APOE and 18F-Florbetapir SUVR**

APOE genotype was significantly associated with 18F-florbetapir SUVR ($\chi^2 = 708.93; P < .001$). Every APOE allele combination was significantly different from all other combinations with the exception of ε22 vs ε23 (1.02 vs 1.02; $P = .91$) and ε22 vs ε33 (1.02 vs 1.05; $P = .43$); note the small sample size of the ε22 group (n = 25). A sample size of 272 for the ε22 group was required to detect the observed difference from the ε33 group with 80% power. Mean 18F-florbetapir SUVR estimates and CIs are summarized in Table 1 and shown in Figure 1. Notably, the ε23 group had a significantly lower mean 18F-florbetapir SUVR compared with the ε33 group (1.02 vs 1.05; $P = .01$), and the ε24 group had a significantly lower mean 18F-florbetapir SUVR compared with the ε34 group (1.11 vs 1.18; $P < .001$). Adjusting for cardiovascular risk score did not affect the APOE genotype estimates and was not associated with 18F-florbetapir SUVR ($\beta = 0.0001; P = .98$).

**APOE, Age, and 18F-Florbetapir SUVR**

There was a significant interaction between APOE genotype and age to predict the odds of Aβ positivity ($\chi^2 = 3.94; P = .41$) (Figure 1).

**Aβ, APOE, and the PACC**

The association between 18F-florbetapir SUVR and decreasing PACC scores did not differ by APOE genotype ($\Delta \text{AIC} = 23.4; P = .97$). Cardiovascular risk was associated with worse PACC scores ($\beta = –0.08; P < .001$) but did not affect the interaction between age and APOE genotype to predict PACC scores ($\Delta \text{AIC} = 22.9; P = .96$). There was also no difference when comparing ε4 carriers to ε4 noncarriers ($\Delta \text{AIC} = 4.4; P = .67$) (Figure 2).

When adjusting for age, sex, and education but not 18F-florbetapir SUVR, the APOE ε4 group performed 0.08 points worse on the PACC compared with the APOE ε33 group ($\beta = 0.08; P = .01$). There were no other significant differences on the PACC compared with the ε33 group ($\beta = –0.25; P = .19$; APOE ε23: $\beta = –0.01; P = .89$; APOE ε24: $\beta = –0.13; P = .14$; APOE ε44: $\beta = –0.11; P = .20$) (Figure 2). Mean PACC scores and 95% CIs for ε22, ε23, ε33, ε34, and ε44 were -0.29 (-0.67 to 0.10); -0.04 (-0.13 to 0.04); -0.04 (-0.08 to 0.00); -0.17 (-0.34 to 0.01); -0.12 (-0.17 to -0.07); and -0.14 (-0.30 to 0.01), respectively (Table 2).

When adjusting for cardiovascular risk, all APOE estimates remained similar, including the effect of APOE ε34 ($\beta = –0.086; P = .007$). When also adjusting for 18F-florbetapir SUVR, the effect of the APOE ε34 group was removed ($\beta = 0.012; P = .71$). For the ε4 carriers vs noncarriers, the unadjusted difference was -0.084 ($P = .005$); after adjusting for 18F-florbetapir, the difference was -0.006 ($P = .85$), and after adjusting for 18F-florbetapir and cardiovascular scores, the difference was -0.009 ($P = .78$).

**Aβ, APOE, and the PACC Components**

The association between 18F-florbetapir SUVR and decreasing PACC component scores did not differ by APOE genotype...
The main findings of this study were (1) APOE ε2 was associated with a reduction in both the overall and the age-dependent level of Aβ in the presence of ε4, (2) large differences in Aβ between APOE groups were already apparent at age 65 years, (3) the association between Aβ and decreasing PACC scores did not differ by APOE genotype, and (4) the associated reduction in performance of the PACC in APOE ε4 carriers compared with noncarriers was completely mediated by Aβ.

There was a large protective outcome of APOE ε2 in the presence of APOE ε4 (Table 1 and Figure 1). β-Amyloid levels in the APOE ε24 group increased at less than half the rate with respect to increasing age compared with the APOE ε34 group. A 2015 meta-analysis did not find a protective effect of carrying the ε2 allele in the presence of ε4 with respect to Aβ positivity.

However, this study had one-third of the number of ε24 participants (n = 41) compared with the A4 Study, most of whom were younger than 70 years. In Figure 1, separation between the ε24 and ε34 groups becomes clear as the groups approach 70 years of age. The reduced levels of Aβ in ε24 compared with ε34 participants shown here may be one of the primary drivers behind the protective outcome of the ε2 allele against AD dementia, shown previously in a large case-control study. The ε24 group demonstrated an associated reduced risk of AD dementia (odds ratio, 2.68 [95% CI, 1.65–4.36]) compared with the ε34 group (odds ratio, 6.13 [95% CI, 5.08–7.41]) when comparing both groups with ε33 participants. However, the presence of the ε2 allele does not completely protect against Aβ positivity, as 16% of ε2 homozygotes in the A4 Study had positive Aβ levels, nor does it completely protect against AD dementia, as 5% of ε24 ε2 homozygotes had a neuropathologically confirmed AD dementia diagnosis. Although the APOE genotype is one of the strongest risk factors for AD, it does not determine Aβ accumulation or cognitive decline.

One of the largest studies to date evaluating age and APOE (including 2914 participants without cognitive impairment) found Aβ positivity in 13.2% of APOE ε4-negative participants and 37.8% of APOE ε4-positive participants at 65 years and 27.7%

### Table 2. Association of Cognition With APOE genotype

<table>
<thead>
<tr>
<th>Measure</th>
<th>ε22 (n = 25)</th>
<th>ε23 (n = 449)</th>
<th>ε33 (n = 4209)</th>
<th>ε24 (n = 115)</th>
<th>ε34 (n = 1295)</th>
<th>ε44 (n = 139)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PACC</td>
<td>-0.29 (-0.67 to 0.10)</td>
<td>-0.04 (-0.13 to 0.04)</td>
<td>-0.04 (-0.08 to 0.00)</td>
<td>-0.17 (-0.34 to 0.01)</td>
<td>-0.12 (-0.17 to -0.07)</td>
<td>-0.14 (-0.30 to 0.01)</td>
</tr>
<tr>
<td>FCSRT96</td>
<td>74.0 (71.8 to 76.3)</td>
<td>77.0 (76.4 to 77.5)</td>
<td>76.5 (76.3 to 76.7)</td>
<td>76.7 (75.7 to 77.7)</td>
<td>76.0 (75.7 to 76.3)</td>
<td>75.3 (74.3 to 76.2)</td>
</tr>
<tr>
<td>Logical Memory Delayed Recall</td>
<td>12.1 (10.8 to 13.4)</td>
<td>11.9 (11.6 to 12.2)</td>
<td>11.8 (11.6 to 11.9)</td>
<td>11.8 (11.2 to 12.3)</td>
<td>11.5 (11.4 to 11.7)</td>
<td>11.9 (11.4 to 12.4)</td>
</tr>
<tr>
<td>Digit Symbol Substitution</td>
<td>45.0 (41.5 to 48.4)</td>
<td>43.1 (42.3 to 43.9)</td>
<td>44.1 (43.8 to 44.4)</td>
<td>41.0 (39.5 to 42.6)</td>
<td>43.8 (43.3 to 44.2)</td>
<td>42.9 (41.5 to 44.3)</td>
</tr>
<tr>
<td>MMSE</td>
<td>28.3 (27.8 to 28.8)</td>
<td>28.8 (28.7 to 28.9)</td>
<td>28.8 (28.6 to 28.9)</td>
<td>28.8 (28.6 to 29.0)</td>
<td>28.8 (28.7 to 28.8)</td>
<td>28.8 (28.6 to 29.0)</td>
</tr>
</tbody>
</table>

Abbreviations: APOE, apolipoprotein E; FCSRT96, Free and Cued Selective Reminding Test; MMSE, Mini-Mental State Examination; PACC, Preclinical Alzheimer Cognitive Composite.

*P > .87; ΔAIC>20.9 for all. When adjusting for demographics but not 18F-florbetapir SUVR or cardiovascular risk, Digit Symbol Substitution Test scores in the APOE ε24 group were reduced (β = -3.06; P = .003). Mean cognitive scores and 95% CIs for ε22, ε23, ε33, ε24, ε34, and ε44 were 45.0 (41.5 to 48.4); 43.1 (42.3 to 43.9); 44.1 (43.8 to 44.4); 41.0 (39.5 to 42.6); 43.8 (43.3 to 44.2); and 42.9 (41.5 to 44.3), respectively (Table 2). This outcome remained after adjusting for 18F-florbetapir SUVR (β = -2.94; P = .005) and also after adjusting for cardiovascular risk (β = -2.97; P = .005).

### Discussion

The main findings of this study were (1) APOE ε2 was associated with a reduction in both the overall and the age-dependent level of Aβ in the presence of ε4, (2) large differences in Aβ between APOE groups were already apparent at age 65 years, (3) the association between Aβ and decreasing PACC scores did not differ by APOE genotype, and (4) the associated reduction in performance of the PACC in APOE ε4 carriers compared with noncarriers was completely mediated by Aβ.

One of the largest studies to date evaluating age and APOE (including 2914 participants without cognitive impairment) found Aβ positivity in 13.2% of APOE ε4-negative participants and 37.8% of APOE ε4-positive participants at 65 years and 27.7%
and 67.8%, respectively, at 80 years. However, a limitation of previous studies is the focus on Aβ positivity (using dichotomous data with conservative thresholds) to define preclinical AD. Using continuous 18F-florbetapir data allows for the estimation of the first increases in Aβ at subthreshold levels. In APOE ε4 carriers, the mean 18F-florbetapir levels were already greatly increased at the minimum age (65 years) compared with APOE ε4 noncarriers. The APOE ε34 and APOE ε44 groups had mean SUVRs of approximately 1.10 and 1.25 at age 65, whereas the ε33 and ε23 groups had mean SUVRs near 1.0. APOE ε4 carrier longitudinal rates of global 18F-florbetapir change have been estimated to be 0.0044 SUVR per year in Aβ-negative individuals and 0.0126 SUVR per year in Aβ-positive individuals, suggesting that it would take decades for the APOE ε4 carrier groups to reach the Aβ levels observed at age 65 years in this study. This coincides with the estimated prevalence of Aβ positivity in ε4 individuals between 25% and 30% at age 45 years and 10% in ε34 individuals at age 50 years. With Aβ positivity already observable in some individuals in their 40s, the gradual accumulation likely begins much earlier. Indeed, reductions of cerebrospinal fluid Aβ have been observed in ε4 carriers in their 20s. A protein-modifying treatment mimicking the protective effect of the ε2 allele against Aβ accumulation may be most effective before significant Aβ deposition. The age at which such a treatment should be initiated would vary greatly by APOE genotype and individual, but if done safely, it could be used to slow Aβ accumulation in early middle age for those at highest risk.

The association between Aβ and performance on the PACC did not differ by APOE genotype. In a recent meta-analysis of 3 large preclinical AD studies, 2 of the 3 studies did not find an interaction between Aβ and APOE genotype to predict cognitive decline. The current study, with a sample size 4 times the size of the previous 3 studies combined, shows the same mean PACC score for a given level of Aβ regardless of APOE genotype. When adjusting for Aβ levels, the significant reduction of PACC scores by 0.08 points in ε4 carriers without Aβ adjustment was completely removed. This reduction is quite modest and refers to a reduction of 0.08 SDs within the Aβ-negative group. However, considering that these participants lack cognitive impairment, this outcome may indicate a nonignorable initial level of cognitive dysfunction. This suggests that the negative association of APOE ε4 positivity on cognition is likely mediated entirely by Aβ accumulation at the preclinical stage of AD. Although many Aβ-independent mechanisms of APOE ε4 have been described, these findings suggest that such Aβ-independent associations of APOE ε4 positivity do not markedly contribute to cognitive decline in the early stages of AD.

Reduced Digit Symbol Substitution test scores in APOE ε2 carriers were unexpected. Although the APOE ε2 allele shows clear protective outcomes against Aβ accumulation, it is also associated with increased risk of atherosclerosis, which in turn is linked to increased risk of cognitive decline and vascular dementia. The associated reduction in executive function and processing speed in the APOE ε24 group observed in this cohort may reflect a non-Alzheimer path to cognitive dysfunction, especially as the adjustment for Aβ burden showed no mediating outcome. Additionally, increased cardiovascular risk scores were associated with decreased cognition in the A4 Study and were highest in the APOE ε24 group, although adjusting for cardiovascular risk factors did not mediate the reduced Digit Symbol Substitution scores observed in the APOE ε24 group. These analyses were exploratory and need to be replicated with longitudinal cognitive trajectories. Still, although the ε2 allele appears to confer protection against Aβ accumulation, safely developing a treatment that mimics its protective outcome will require care not to increase cardiovascular risk or another non-Aβ path to cognitive dysfunction.

Limitations
This study has several limitations. This study was limited to participants without cognitive impairment older than 65 years of age, thereby limiting analyses to the outcome of emerging Aβ pathology in the absence of significant cognitive dysfunction. Importantly, this is a cross-sectional study, and these results do not apply to changes within individuals. Longitudinal follow-up of early middle-aged individuals, with low and intermediate levels of amyloid, will be required to further clarify when APOE groups initially diverge. Importantly, participation in an AD prevention trial is voluntary, which may introduce bias and reduce generalizability. A4 Study participants are highly educated relative to the general population. The vast majority of A4 Study participants are White and are not representative of the population at risk for AD. Exclusion criteria limited participation to those without health conditions that could interfere with the study, which may introduce bias. Although the sample size of our primary group of interest, ε24 carriers, was relatively large compared with previous studies, further subdivision by factors known to be associated with APOE genotype, particularly race/ethnicity, were precluded by small sample sizes. Finally, use of the PACC, a cognitive composite, may be limited in its sensitivity to specific cognitive differences depending on the distribution across cognitive domains.

Conclusions
This study’s findings suggest that the protective outcome of carrying an ε2 allele in the presence of an ε4 allele offers potential for a treatment that attempts to mimic this protective outcome in order to facilitate Aβ clearance in ε4 carriers. Such a treatment strategy is appealing, as ε4 carriers make up 67% of patients with AD dementia, and it could represent an early treatment option, as many ε4 carriers begin to accumulate Aβ in early middle age. If the goal is to interfere early in the disease process before activation of downstream pathways, AD prevention trials may consider targeting much younger people before the accumulation of high or even intermediate levels of Aβ develop.
Apolipoprotein E ε2 vs ε4, Age, and β-Amyloid in Adults Without Cognitive Impairment

Drafting of the manuscript: Insel, Mattsson-Carlgren.

Critical revision of the manuscript for important intellectual content: All authors.

Statistical analysis: Insel.

Obtained funding: Mattsson-Carlgren.

Supervision: Hansson, Mattsson-Carlgren.

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Additional Information: The Anti-Amyloid Treatment in Asymptomatic Alzheimer Disease Study (A4 Study) is a secondary prevention trial in preclinical Alzheimer disease, aiming to slow cognitive decline associated with brain amyloid accumulation in clinically normal older individuals. The A4 Study is funded by a public-private-philanthropic partnership, including funding from the National Institutes of Health-National Institute on Aging (U01AG010483; R01AG063689), Eli Lilly and Company, Alzheimer Association, Accelerating Medicines Partnership, GHR Foundation, an anonymous foundation, and additional private donors, with in-kind support from Avid, Cogstate, Albert Einstein College of Medicine, US Against Alzheimer Disease, and Foundation for Neurologic Diseases. The companion observational Longitudinal Evaluation of Amyloid Risk and Neurodegeneration Study is funded by the Alzheimer Association and GHR Foundation. The A4 and Longitudinal Evaluation of Amyloid Risk and Neurodegeneration Studies are led by Reisa Sperling, MD, at the Brigham and Women's Hospital, Harvard Medical School and Paul Aisen, MD, at the Alzheimer Therapeutic Research Institute, University of Southern California. The A4 and Longitudinal Evaluation of Amyloid Risk and Neurodegeneration Studies are coordinated by the Alzheimer Therapeutic Research Institute at the University of Southern California, and the data are made available through the Laboratory for Neuro Imaging at the University of Southern California. The participants screening for the A4 Study provided permission to share their deidentified data in order to advance the quest to find a successful treatment for Alzheimer disease. The complete A4 Study Team list is available at a4study.org/a4-study-team.

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REFERENCES


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