Comparing cortical signatures of atrophy between late-onset and autosomal dominant Alzheimer disease

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Abbreviations: AD, Alzheimer disease; ADAD, autosomal dominant Alzheimer disease; APOE, apolipoprotein E; APP, amyloid precursor protein; AUROC, area under the receiver operating characteristic; AV-45, florbetapir; CDR, clinical dementia rating; CN_ADRD/DIAN, cognitively normal controls; DIAN, Dominantly Inherited Alzheimer Network; HCV, total hippocampal volume; Knight ADRC, Knight Alzheimer Disease Research Center; LOAD, late-onset Alzheimer disease; mcSUVR, mean cortical standardized uptake value ratio; MRI, magnetic resonance imaging; PCalcld, preclinical Alzheimer disease; P_CG3, preclinical autosomal dominant Alzheimer disease; PET, positron emission tomography; PiB, Pittsburgh compound-B; PSEN1, Presenilin 1; PSEN2, Presenilin 2; ROC, receiver operating characteristic; ROI, region of interest; SUVR, standardized uptake value ratio.

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ABSTRACT
Defining a signature of cortical regions of interest preferentially affected by Alzheimer disease (AD) pathology may offer improved sensitivity to early AD compared to hippocampal volume or mesial temporal lobe alone. Since late-onset Alzheimer disease (LOAD) participants tend to have age-related comorbidities, the younger-onset age in autosomal dominant AD (ADAD) may provide a more idealized model of cortical thinning in AD. To test this, the goals of this study were to compare the degree of overlap between the ADAD and LOAD cortical thinning maps and to evaluate the ability of the ADAD cortical signature regions to predict early pathological changes in cognitively normal individuals.

We defined and analyzed the LOAD cortical maps of cortical thickness in 588 participants from the Knight Alzheimer Disease Research Center (Knight ADRC) and the ADAD cortical maps in 269 participants from the Dominantly Inherited Alzheimer Network (DIAN) observational study. Both cohorts were divided into three groups: cognitively normal controls (n_{ADRC} = 381; n_{DIAN} = 145), preclinical (n_{ADRC} = 153; n_{DIAN} = 76), and cognitively impaired (n_{ADRC} = 54; n_{DIAN} = 48). Both cohorts underwent clinical assessments, 3T MRI, and amyloid PET imaging with either 11C-Pittsburgh compound B or 18F-florbetapir.

To generate cortical signature maps of cortical thickness, we performed a vertex-wise analysis between the cognitively normal controls and impaired groups within each cohort using six increasingly conservative statistical thresholds to determine significance. The optimal cortical map among the six statistical thresholds was determined from a receiver operating characteristic analysis testing the performance of each map in discriminating between the cognitively normal controls and preclinical groups. We then performed within-cohort and cross-cohort (e.g., ADAD maps evaluated in the Knight ADRC cohort) analyses to examine the sensitivity of the optimal cortical signature maps to the amyloid levels using only the cognitively normal individuals (cognitively normal controls and preclinical groups) in comparison to hippocampal volume.

We found the optimal cortical signature maps were sensitive to early increases in amyloid for the asymptomatic individuals within their respective cohorts and were significant beyond the inclusion of hippocampal volume, but the cortical signature maps performed poorly when analyzing across cohorts. These results suggest the cortical signature maps are a useful MRI biomarker of early AD-related neurodegeneration in preclinical individuals and the pattern of decline differs between LOAD and ADAD.

1. Introduction
Alzheimer disease (AD) is defined pathologically by the accumulation of amyloid plaques and neurofibrillary tangles. Amyloid pathology is detectable almost 20 years before symptom onset and is followed by limbic and neocortical neurofibrillary tangles (Bateman et al., 2012; Hardy and Higgins, 1992; Jack et al., 2013). Cognitively, AD manifests as a cortical dementia. Cerebral atrophy occurs nearer to onset and is more closely related, spatially and temporally, to cognitive decline. (Aschenbrenner et al., 2018; Brier et al., 2016a; Ossenkoppele et al., 2019). Estimates of volume and cortical thickness from magnetic resonance imaging (MRI) are one of the most common measures of neurodegeneration and have been incorporated into recommendations for using biomarkers to study AD (Jack et al., 2018). Determining the specific brain regions most susceptible to early neurodegeneration in AD would help identify individuals most at risk of cognitive decline.

Guided by neuropathological studies (Braak and Braak, 1995), much of AD research has focused on atrophy of the mesial temporal lobe and the hippocampus in particular (Apostolova et al., 2010; Bouwman et al., 2007; Convit et al., 1997; Gordon et al., 2016; Huijbers et al., 2015; Martin et al., 2010; Ridha et al., 2006; Vos et al., 2016). Late-onset Alzheimer disease (LOAD) is the most common form of AD and typically develops in individuals aged 65 and older. However, older healthy and AD individuals are known to have a high frequency of comorbidities, which can affect brain volumetrics (e.g., diabetes, hypertension, cerebrovascular disease) including hippocampal size (e.g., hippocampal sclerosis) (Faraco and Iadecola, 2013; Jagust et al., 2008; Moran et al., 2013; Zarow et al., 2008). These comorbidities make the hippocampus a relatively nonspecific structure for measuring neurodegeneration.

Looking beyond the hippocampus, previous studies have defined a composite of brain regions susceptible to AD-related cortical atrophy, known as AD cortical signatures (Dickerson et al., 2009; Jack et al., 2017; Wang et al., 2015) and have shown cortical signatures relate to cognitive performance in AD beyond the effects of age (Bakkour et al., 2013). Voxel or vertex-wise approaches (Dickerson et al., 2009; Wang et al., 2015) with modest sample sizes often led to statistical maps with a series of peak effects spread throughout cortical and subcortical areas. Approaches using a larger region of interest (ROI) derived from programs such as FreeSurfer avoid the small focal effects but may incorporate areas of the brain not affected by AD. We therefore propose that an ideal signature region should be large enough to be stable across multiple cohorts, yet specific enough that it does not incorporate non-informative regions.

Although AD cortical signatures include numerous brain regions, the age-related comorbidities in the LOAD population may still limit the sensitivity to detect AD-related neurodegeneration in asymptomatic individuals. Autosomal dominant Alzheimer disease (ADAD) is a rare form of AD caused by a mutation in one of three genes: amyloid precursor protein (APP), presenilin1 (PSEN1), or presenilin2 (PSEN2). ADAD has virtually complete penetrance and a consistent age of onset within families for each individual mutation (Bateman et al., 2010). Since ADAD has an earlier age of onset compared to LOAD (Ryman et al., 2014), ADAD individuals typically do not have age-related comorbidities. Prior work looking at atrophy in ADAD cohorts using ROI (Benzinger et al., 2013; Gordon et al., 2018a; Weston et al., 2016) and voxel-wise approaches (Scähill et al., 2013, 2002) found a robust atrophy signature. Therefore, ADAD may provide a less biased depiction needed to identify changes in the preclinical stages of AD.

In this study, we sought to produce a refined AD cortical atrophy signature by 1) creating a signature without ROI restrictions and with a large cohort for more reliable regions, 2) comparing the degree of overlap between the ADAD and LOAD signatures, and 3) comparing the abilities of the ADAD and LOAD signatures to predict early amyloid pathological changes in the cognitively normal groups within and across cohorts.
2. Materials and methods

2.1. Participants

Two datasets were used to define and analyze AD cortical signatures of cortical thickness. For both datasets, dementia severity was defined by the global Clinical Dementia Rating (CDR) (Morris, 1993). The LOAD participants were selected from ongoing longitudinal studies of memory and aging at the Knight Alzheimer Disease Research Center (Knight ADRC) at Washington University in Saint Louis. All participants from the 12th semiannual Knight ADRC data release that had at least one 3 Tesla (3T) MRI scan, an amyloid positron emission tomography (PET) scan within a year of the MRI, a clinical assessment within two years of the MRI, and demographic data including age, sex, and apolipoprotein E (APOE) genotype were included in the study (n = 703). The date of the MRI scan was considered the baseline visit. Participants were subsequently divided into three groups: cognitively normal controls (CNLOAD, CDR 0 and amyloid negative for all visits, n = 381), preclinical LOAD (PCLOAD, CDR 0 and amyloid positive at baseline visit, n = 153), and cognitively impaired (LOAD, CDR 0.5 and amyloid positive at baseline visit and remains CDR 0.5 or higher at subsequent visits, n = 54). 115 participants were excluded as follows: 43 participants were CDR 0.5 and amyloid negative; 37 participants were CDR 0 and amyloid negative at baseline visit but did not remain so for all other visits; 26 participants were CDR > 0.5 at baseline visit; 6 participants were CDR 0.5 and amyloid positive at baseline visit but did not remain CDR 0.5 or higher for subsequent visits; 3 participants had a non-AD diagnosis.

ADAD participants were selected from the Dominantly Inherited Alzheimer Network (DIAN) observational study (Morris et al., 2012; Moulder et al., 2013). Participants are recruited from families having an ADAD pathogenic mutation in APP, PSEN1, or PSEN2 genes. Participant enrollment and procedures have been previously defined (Morris et al., 2012). For this study, participants from the 11th DIAN annual data release were required to have the same criteria as the Knight ADRC cohort (n = 386). Participants were then divided into three groups: cognitively normal mutation-non-carrier controls (CNADAD, CDR 0 and amyloid negative for all time points, n = 145), preclinical ADAD mutation carriers (PCADAD, CDR 0 and amyloid positive at baseline visit, n = 76), and cognitively impaired mutation carriers (ADAD, CDR 0.5 and amyloid positive at baseline visit and remains CDR 0.5 or higher at subsequent visits, n = 48). 117 participants were excluded as follows: 65 participants were CDR 0 and amyloid negative mutation carriers; 29 participants were CDR > 0.5 at baseline visit; 18 participants were CDR 0.5 and amyloid negative; 2 participants were CDR 0 and amyloid negative mutation non-carriers at baseline visit but did not remain so for all other visits; 2 participants were CDR 0 and amyloid positive mutation non-carriers; 1 participant was CDR 0.5 and an amyloid positive mutation carrier but did not remain CDR 0.5 or higher for subsequent visits.

2.2. Ethics statement

All participants or their legal caregivers provided written informed consent approved by their local institution’s review board. Each site’s institutional review board approved all study procedures. For the DIAN study, the institutional review board at Washington University in St. Louis provided supervisory review and human studies approval.

2.3. MRI acquisition and processing

For the Knight ADRC cohort, T1-weighted images were acquired using a magnetization-prepared rapid gradient-echo sequence on either the Siemens Biograph mMR (n = 172) or the Siemens 3T TIM Trio (n = 416) scanner. Structural T1 scans for the Biograph mMR were acquired with a 1 × 1 × 1.2 mm resolution, 2300 ms repetition time, 2.95 ms echo time, 9 degree flip angle, 176 frames, and a 240 × 256 field of view in sagittal orientation. Structural T1 scans for the TIM Trio were acquired with a 1 × 1 × 1 mm resolution, 2400 ms repetition time, 3.16 echo time, 8 degree flip angle, 176 frames, and a 256 × 256 field of view in sagittal orientation. For the DIAN cohort, the T1-weighted images were acquired using the Alzheimer Disease Neuroimaging Initiative structural MRI protocol (Jack et al., 2010) using either a Siemens (n = 236) or a Phillips (n = 33) 3T scanner and the images underwent quality control assessments for protocol compliance. Structural T1 scans had a resolution of 1 × 1 × 1.2 mm. Images from both cohorts were processed with FreeSurfer 5.3-HCP (Fischl et al., 2002) and were resampled to 1 × 1 × 1 mm resolution for volumetric segmentation and cortical reconstruction. The total hippocampal volume (HCV) was derived from the sum of the left and right hemisphere hippocampal volume from the automated FreeSurfer regions.

2.4. PET acquisition and processing

Amyloid PET imaging was performed with 11C-Pittsburgh compound B (PiB) in the DIAN cohort and either PiB or 18F-florbetapir (AV-45) in the Knight ADRC cohort. All PET scans were processed with the PET Unified Pipeline (PUP, https://github.com/ysu001/PUP) using FreeSurfer derived ROIs (Su et al., 2015, 2013) to calculate the standardized uptake value ratio (SUVR) using the cerebellar cortex as a reference region. A 30–60 min post-injection window for PiB and a 50–70 min post-injection window for AV-45 were used in the Knight ADRC cohort and a 40–70 min post injection window for PiB was used in the DIAN cohort. Partial volume correction was performed using a geometric transfer matrix (Rousset et al., 1998; Su et al., 2015). To measure amyloid burden, we calculated the mean cortical SUVR (mcSUVR) by averaging the partial volume corrected SURVs from the FreeSurfer ROIs in the lateral orbitofrontal, mesial orbitofrontal, rostral mesial frontal, superior frontal, superior temporal, mesial temporal, and precuneous regions as previously defined (Su et al., 2013).

To standardize the PiB and AV-45 tracers, the mcSUVR values for the Knight ADRC cohort were converted into Centiloid units (Klunk et al., 2015) using a linear transformation from previously defined equations (Su et al., 2019, 2018). The value of zero on the Centiloid scale represents the mean amyloid burden for healthy participants and the value of 100 on the Centiloid scale represents the mean amyloid burden for AD participants. For statistical analysis, we utilized Centiloid for the Knight ADRC cohort and mcSUVR for the DIAN cohort. Amyloid positivity was defined as an mcSUVR ≥ 1.42 for PiB scans (Brier et al., 2016b; Sutphen et al., 2015) and an mcSUVR ≥ 1.22 for AV-45 scans (Mishra et al., 2017).

2.5. Creation of the cortical maps

For each hemisphere, cortical thickness spatial maps were created via vertex-wise analyses using a general linear model in FreeSurfer. The LOAD cortical maps were generated by computing cortical thickness differences between the LOAD and CNLOAD groups while adjusting for age and sex. Similarly, the ADAD cortical maps were generated by calculating cortical thickness differences between the ADAD and CNDIAN groups while accounting for the effects of age and sex.

Generating a robust cortical signature requires a balance between specificity and sensitivity; it is a tradeoff between having a large enough cortical map which adequately captures the most informative regions but not so large it incorporates regional information with limited predictive value. For this reason, Monte Carlo simulations were performed using a selection of vertex-wise thresholds (p < 0.05, p < 0.01, p < 0.005, p < 0.001, p < 0.0005 and p < 0.0001) followed by a p < 0.001 cluster-wise threshold to correct for multiple comparisons. Each Monte Carlo vertex-wise threshold generated a different ROI, which is identified as the significant cluster of vertices, and were generated separately for the Knight ADRC and DIAN cohorts (Supplementary Fig. 1, Supplementary Fig. 2). The LOAD and ADAD ROIs were mapped to participant space within and across cohorts. For each hemisphere, the mean cortical
thickness values were obtained for each cortical map (Supplementary Fig. 3).

To identify the ROI cortical maps which represent the optimal “cortical signature”, we compared the ability of the mean cortical thickness in each ROI to discriminate between the CN and PC groups by calculating the area under the receiver operating characteristic (AUROC) curve using the pROC package (Robin et al., 2011) in R. The left and right hemisphere ROIs which exhibited the best performance (i.e. those with the highest AUROC values) were identified as the optimal cortical signature and defined separately for the Knight ADRC and DIAN cohorts. Subsequently, the thickness values from the optimal cortical maps were averaged across hemispheres and weighted by the number of vertices in the ROI to obtain the cortical signature thickness. The optimal ADAD and LOAD cortical signatures are available online at https://github.com/benzinger-icl/ADcortsig-roi.

To verify that the cortical signature thickness performed with similar results to the receiver operating characteristic (ROC) analysis, a within-cohort analysis was done examining the cortical signature thickness performance from the ADAD and LOAD maps in predicting PC versus CN, and ADAD versus CNAD, respectively. Second, we performed a cross-cohort analysis examining the cortical signature thickness performance from the ADAD and LOAD maps in predicting PCDIAN versus CNAD, and PDIVAN versus CNDIVAN, respectively. As atrophy is a biomarker that occurs before but close to symptom onset, we also repeated the cross-cohort analysis in a subset of the PCDIAN (n = 39) and PDIVAN (n = 19) participants who were in the top quartile in terms of amyloid burden.

2.6. Statistical analysis

All statistical tests were conducted in R version 3.6.0 (R Core Team, 2019). Logistic regression models were used to examine the ability of cortical signature thickness to discriminate PC versus CN groups with amyloid status as the dependent term and the cortical signature thickness as the independent term while adjusting for age and sex. Additionally, we performed linear regression models to examine the ability of cortical signature thickness to predict amyloid burden in the PC and CN groups. The amyloid burden measure was the dependent term, defined in Centiloid units for the Knight ADRC cohort and mcSUVR for the DIAN cohort, while the cortical signature thickness measure was the independent term with adjustments made to account for age and sex variability. The purpose of the linear models is to examine the amyloid levels immediately above and below the amyloid status cutoff as these values are meaningful and may provide a more sensitive measure to detect the PC and CN groups. The logistic and linear models were again run including HCV as a covariate as a direct comparison of MRI biomarkers. Due to the prevalence of comorbidities in LOAD, we additionally examined the association of obesity, measured with body mass index (BMI), and white matter hyperintensity (WMH) burden, measured with the log transformed WMH volume, against the LOAD cortical signature thickness for a subset of the CNAD and PCDIAN groups (nWMH = 237) by calculating the Pearson correlation coefficient (r). Finally, we were interested in quantifying demographic differences within each cohort. For our continuous variables, age and education, ANOVA were run and follow-up pair-wise t-tests were conducted for any significant outcomes. Similarly, for our categorical variables, APOE ε4 status and sex, chi-square tests were run and follow-up pair-wise chi-square tests were conducted for significant outcomes.

3. Results

3.1. Demographics

The demographics from the DIAN and Knight ADRC cohorts are presented in Table 1.

Table 1

| Table 1 | DIAN Cohort | Knight ADRC Cohort |
|---------|-------------|-------------------|
|         | CN_DIAN     | PCDIAN            | ADAD | CN_ADRC | PCDAD | LOAD |
| N       | 145         | 76                | 48   | 381      | 153   | 54   |
| Age, years | 38.5       | 37.2              | 44.8 | 65.5     | 73.1  | 76.5 |
| (11.3)  | (9.0)       | (10.6)            | (9.4) | (7.1)    | (7.1)     |
| EYO     | –9.9        | –9.9              | 3.0  | –        | –     | –    |
| (11.8)  | (7.8)       | (2.7)             | –    | –        | –     | –    |
| Male, n (%) | 58 (17)    | 58 (17)            | 159  | 66 (11.8) | 15.1  | 29   |
| (0.0)  | (0.0)       | (48.7)            | (37.5) | (43.1)  | (53.7) |
| Education, years | 15.0   | 14.7              | 13.4 | 16.0     | 16.0  | 15.1 |
| (2.7)  | (2.9)       | (2.5)             | (2.4) | (2.4)    | (2.4) |
| APOE ε4, n (%) | 42 (26)  | 21 (15)            | 106  | 88 (41)  | 44    | 37   |
| (20.0) | (20.0)      | (34.2)            | (31.2) | (37.8)  | (37.8)|
| PSEN1/ PSEN2/APP, n | 97 / 18 | 97 / 30             | 59 (75.9) | 39 / 2   | 106  | 145  |
| / 30 / 6 / 7 | –       | –                 | –    | –        | –     | –    |
| CDR 0/0.5, n | 145 / 0 | 76 / 0             | 48   | 0 / 0    | 0 / 0 | 54   |
| 0.0     | 0 / 48      | 381 / 0            | 153 / 0 | 0 / 0    | 0 / 0 |
| Aβ1-42/Aβ1-42, n | 145 / 0 | 76 / 0             | 48   | 0 / 0    | 0 / 0 | 54   |
| 0.0     | 0 / 48      | 381 / 0            | 153 / 0 | 0 / 0    | 0 / 0 |
| ADAD cortical signature thickness, mm | 2.34 (0.11) | 2.31 (0.13)  | 2.09 (0.17) | 2.22 (0.12) | 2.16 (0.12) | 2.08 (0.12) |
| LOAD cortical signature thickness, mm | 2.74 (0.12) | 2.73 (0.12)  | 2.60 (0.15) | 2.60 (0.12) | 2.53 (0.12) | 2.36 (0.12) |
| PIB-PET SUVR | 1.05 (0.07) | 2.05 (0.61)  | 2.89 (1.08) | 0.1 (48.2) | 82.5 | 82.5 |
| Centiloid | –0.4 (3.1) | 44.8 (27.5)  | 82.5 (48.6) | 6.2 (62.6) | 26.0 (26.7) |

Mean (standard deviation) unless otherwise noted.

ADAD = autosomal dominant Alzheimer disease; LOAD = late-onset Alzheimer disease; CN_DIAN = cognitively normal mutation non-carrier; CN_ADRC = cognitively normal controls; PCDIAN = preclinical ADAD; PCDAD = preclinical LOAD; EYO = estimated years to symptom onset; APOE = apolipoprotein E; PSEN1 = presenilin 1; PSEN2 = presenilin 2; APP = amyloid precursor protein; CDR = Clinical Dementia Rating; Aβ1-42/Aβ1-42 = amyloid negative/amyloid positive; PIB-PET = Pittsburgh compound B positron emission tomography.

a Significant differences between CN and PC, p < 0.05.
b Significant differences between CN and ADAD/LOAD, p < 0.05.
c Significant differences between PC and ADAD/LOAD, p < 0.05.

were found between the CN_DIAN versus ADAD groups and PCDIAN versus ADAD groups. However, there were no significant results for sex or APOE ε4 status from these analyses. In contrast, we report significant age and APOE ε4 status differences for all groups within the Knight ADRC cohort, but there were no significant differences in education and sex for these groups.

3.2. ADAD and LOAD cortical signatures

The ADAD cortical maps for each vertex-wise threshold are presented in Supplementary Fig. 1. The AUROC values from the ROC analysis discriminating PCDIAN and CN_DIAN groups for each hemisphere and each vertex-wise significance threshold (p < 0.05, p < 0.01, p < 0.005, p < 0.001, p < 0.0005, p < 0.0001) are presented in Table 2.

The highest AUROC value for the right hemisphere was 0.5469, which corresponded to the cortical map generated using the p < 0.001 threshold. For the left hemisphere, the highest AUROC value was 0.5469, which corresponded to the cortical map defined using the p < 0.0001 threshold. These two cortical maps define the optimal cortical signature for ADAD shown in Fig. 1A. The associated effect size maps are presented in Fig. 1B. We also identified the approximate FreeSurfer regions which best match the vertex-wise signature by calculating the percentage of voxels in the signature map overlaying each FreeSurfer region. Specifically, the vertex-wise signature was converted into a volume and the percentage of
overlapping voxels was determined by dividing the number of cortical signature voxels within a FreeSurfer region by the total number of voxels in that FreeSurfer region and multiplying by 100. The FreeSurfer region with the highest ADAD signature overlap for the right hemisphere is the inferior parietal region (60% overlap) and for the left hemisphere is the precuneus region (54% overlap).

Identical analyses were performed in the Knight ADRC cohort to determine the optimal cortical signature. The LOAD cortical maps for all vertex-wise thresholds are presented in Supplementary Fig. 2 and the AUROC values for each hemisphere and threshold are presented in Table 2. The best AUROC was 0.6470 for the right hemisphere, corresponding to the \( p < 0.05 \) vertex-wise threshold, and 0.6518 for the left hemisphere, corresponding to the \( p < 0.005 \) threshold. These two cortical maps are used to define the optimal LOAD cortical signature presented in Fig. 2A. The associated effect size maps of the cortical signature are found in Fig. 2B.

The FreeSurfer region with the greatest LOAD signature overlap for the left and right hemisphere is the entorhinal cortex (85% right hemisphere overlap and 74% left hemisphere overlap). A list of additional FreeSurfer regions and the associated cortical signature overlap percentage are presented in Supplemental Table 1. We additionally compared the ADAD and LOAD cortical signatures (Fig. 3). The LOAD signature emphasized atrophy in the temporal lobe whereas ADAD is more prominent in the parietal and precuneus. There is also a noticeable overlap within the precuneus and parietal regions in both signatures.

### 3.3. Within-cohort analysis: association of the cortical signature thickness with amyloid

Although each hemispheric signature was optimized to discriminate between PC and CN groups, we wanted to confirm the cortical signature thickness differences between the ADAD and \( \text{CN}_{\text{DIAN}} \) groups including age and sex as covariates and using a \( p < 0.001 \) cluster-wise threshold. The vertex-wise threshold was found to be \( p < 0.001 \) for right hemisphere and \( p < 0.0001 \) for left hemisphere. Significant vertices are depicted in yellow highlighting the inferior parietal and precuneus. B. The cortical thickness effect size maps comparing ADAD and \( \text{CN}_{\text{DIAN}} \). The blue indicates areas where the ADAD group had lower cortical thickness values compared to the \( \text{CN}_{\text{DIAN}} \) and the red indicates areas where the ADAD group had higher cortical thickness values compared to the \( \text{CN}_{\text{DIAN}} \) group. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

**Fig. 1.** A. The optimal ADAD cortical signature depicting cortical thickness differences between the ADAD and \( \text{CN}_{\text{DIAN}} \) groups including age and sex as covariates and using a \( p < 0.001 \) cluster-wise threshold. The vertex-wise threshold was found to be \( p < 0.001 \) for right hemisphere and \( p < 0.0001 \) for left hemisphere. Significant vertices are depicted in blue highlighting predominantly the temporal lobe. B. The cortical thickness effect size maps comparing ADAD and \( \text{CN}_{\text{DIAN}} \). The blue indicates areas where the ADAD group had lower cortical thickness values compared to the \( \text{CN}_{\text{DIAN}} \) and the red indicates areas where the ADAD group had higher cortical thickness values compared to the \( \text{CN}_{\text{DIAN}} \) group. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

**Fig. 2.** A. The optimal LOAD cortical signature depicting cortical thickness differences between the LOAD and \( \text{CN}_{\text{ADRC}} \) groups including age and sex as covariates and using a \( p < 0.001 \) cluster-wise threshold. The vertex-wise threshold was found to be \( p < 0.05 \) for right hemisphere and \( p < 0.005 \) for left hemisphere. Significant vertices are depicted in blue highlighting predominantly the temporal lobe. B. The cortical thickness effect size maps comparing LOAD and \( \text{CN}_{\text{ADRC}} \). The blue indicates areas where the LOAD group had lower cortical thickness values compared to the \( \text{CN}_{\text{ADRC}} \) and the red indicates areas where the LOAD group had higher cortical thickness values compared to the \( \text{CN}_{\text{ADRC}} \) group. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

| Monte Carlo Vertex-wise Thresholds | ADAD ROI Cortical Maps | LOAD ROI Cortical Maps |
|----------------------------------|------------------------|------------------------|
|                                  | RH | LH | RH | LH | RH | LH |
| \( p < 0.05 \)                   | 0.4671 | 0.5146 | 0.6470 | 0.6455 |
| \( p < 0.01 \)                   | 0.5357 | 0.5397 | 0.6332 | 0.6487 |
| \( p < 0.005 \)                  | 0.5330 | 0.5431 | 0.6427 | 0.6518 |
| \( p < 0.001 \)                  | 0.5400 | 0.5329 | 0.6426 | 0.6443 |
| \( p < 0.0005 \)                 | 0.4601 | 0.5396 | 0.6422 | 0.6509 |
| \( p < 0.0001 \)                 | 0.4540 | 0.5469 | 0.6326 | 0.6379 |

**Bold** indicates the highest AUROC value for each ROI and hemisphere.

ADAD = autosomal dominant Alzheimer disease; LOAD = late-onset Alzheimer disease; AUROC = area under the receiver operating characteristic.
Fig. 3. A comparison of the ADAD and LOAD optimal cortical signatures. Yellow indicates ADAD areas, blue indicates LOAD areas, and green indicates areas where both ADAD and LOAD cortical signatures overlap. Much of the overlap between the two signatures are in parts of the precuneus and parietal regions. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

signatures’ capabilities within its own cohort and compare it to a more common MRI biomarker, HCV. We found that the ADAD cortical signature thickness marginally separated \( CN_{\text{DIAN}} \) and \( PC_{\text{DIAN}} \) (ADAD cortical signature thickness: \( \beta = -2.44, p = 0.048 \)). While the ADAD cortical signature thickness discriminated \( CN_{\text{DIAN}} \) and \( PC_{\text{DIAN}} \), the HCV did not reach statistical significance. (ADAD cortical signature thickness: \( \beta = -2.51, p = 0.044; \) HCV: \( \beta = 7.37e-5, p = 0.69 \)). In the linear regression analysis, the ADAD cortical signature thickness predicted amyloid burden within the combined \( PC_{\text{DIAN}} \) mutation carrier and \( CN_{\text{DIAN}} \) mutation non-carrier groups (ADAD cortical signature thickness: \( \beta = -0.78, p = 0.025 \)) whereas HCV was not able to discriminate groups (ADAD cortical signature thickness: \( \beta = -0.76, p = 0.03; \) HCV: \( \beta = -1.42e-5, p = 0.79 \)).

For the Knight ADRC, the LOAD cortical signature thickness did not significantly separate the \( CN_{\text{ADRC}} \) and \( PC_{\text{ADRC}} \) groups (LOAD cortical signature thickness: \( \beta = -0.89, p = 0.36 \)), primarily due to the significant effect of age in the model (age: \( \beta = 0.09, p = 3.74e-10 \)), and HCV performed similarly to the cortical signature (LOAD cortical signature thickness: \( \beta = -0.85, p = 0.41; \) HCV: \( \beta = -1.60e-5, p = 0.90 \)). When treating amyloid as a continuous measure, the LOAD cortical signature thickness predicted amyloid burden (LOAD cortical signature thickness: \( \beta = -24.37, p = 0.016 \)) beyond the covariates but HCV did not reach statistical significance (LOAD cortical signature thickness: \( \beta = -22.14, p = 0.036; \) HCV: \( \beta = -9.70e-4, p = 0.46 \)). Amyloid as a continuous variable performed better in all statistical models when compared to analyzing amyloid as a categorical variable. The distributions for the cortical signature thickness and HCV within the DIAN and Knight ADRC cohorts are shown in Fig. 4.

When analyzing BMI and WMH volume against the LOAD cortical signature thickness in the \( CN_{\text{ADRC}} \) and \( PC_{\text{ADRC}} \) groups, no association was present between BMI and LOAD cortical signature thickness \((r = -0.06).\) Notably, there was an association between the log transformed WMH volume and LOAD cortical signature thickness \((r = -0.63),\) primarily driven by age (Supplementary Fig. 4).

3.4. Cross-cohort analysis: association of the cortical signature thickness with amyloid

Since there are known similarities between ADAD and LOAD, we examined whether the ADAD signature could differentiate between the \( PC_{\text{ADRC}} \) and \( CN_{\text{ADRC}} \) groups and whether the LOAD signature could differentiate between the \( PC_{\text{DIAN}} \) and \( CN_{\text{DIAN}} \) groups. We found the ADAD cortical signature thickness was unable to discriminate the \( PC_{\text{ADRC}} \) and \( CN_{\text{ADRC}} \) groups (ADAD cortical signature thickness: \( \beta = -0.61, p = 0.52 \)) and likewise when including HCV into the model (ADAD cortical signature thickness: \( \beta = -0.55, p = 0.57; \) HCV: \( \beta = -3.47e-5, p = 0.79 \)). The ADAD cortical signature thickness could not predict amyloid burden (ADAD cortical signature thickness: \( \beta = -12.46, \)

Fig. 4. The distributions of the ADAD and LOAD cortical signature thickness and hippocampal volume within the DIAN and Knight ADRC cohorts, respectively. For comparison, the cortical signature thickness and hippocampal volume were normalized relative to the cognitively normal control groups. The black circles indicate the mean for each group. \( CN_{\text{DIAN}} \) = cognitively normal mutation non-carrier (dark green); \( PC_{\text{DIAN}} \) = preclinical ADAD (orange); ADAD = autosomal dominant Alzheimer disease (purple); \( CN_{\text{ADRC}} \) = cognitively normal controls (pink); \( PC_{\text{ADRC}} \) = preclinical LOAD (green); LOAD = late-onset Alzheimer disease (yellow). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
cortical signature thickness: the logistic regression analysis was repeated to only include the top quartile amyloid burden PC participants. In this analysis, the ADAD cortical signature thickness could not discriminate the groups (ADAD cortical signature thickness: \(\beta = -2.08, p = 0.18\)) and was unable to predict amyloid burden (ADAD cortical signature thickness: \(\beta = -16.13, p = 0.14\)).

The LOAD cortical signature thickness was unable to distinguish between the CN\textsubscript{DIAN} and PC\textsubscript{DIAN} groups (LOAD cortical signature thickness: \(\beta = -1.01, p = 0.42\)) and HCV performed similarly (LOAD cortical signature thickness: \(\beta = -1.10, p = 0.39\); HCV: \(\beta = 5.56e-5, p = 0.77\)). Additionally, the LOAD cortical signature thickness could not predict amyloid burden (LOAD cortical signature thickness: \(\beta = -0.38, p = 0.29\)) and likewise for HCV (LOAD cortical signature thickness: \(\beta = -0.35, p = 0.34\); HCV: \(\beta = -1.92e-5, p = 0.72\)). When restricting the PC\textsubscript{DIAN} group to the top quartile amyloid burden participants, the LOAD cortical signature thickness was unable to separate the groups (LOAD cortical signature thickness: \(\beta = -1.86, p = 0.38\)) or predict amyloid burden (LOAD cortical signature thickness: \(\beta = -0.57, p = 0.19\)).

4. Discussion

AD-related atrophy affects many brain regions beyond just the hippocampus alone. LOAD individuals are also likely to have age-related comorbidities which could make it difficult to characterize regions more prone to AD-specific pathology. Since individuals affected by ADAD tend to be younger and thus lack many age-related comorbidities, this cohort may provide a better cortical atrophy model specific to AD. The goals of this study were to create a refined and reliable cortical atrophy signature by using a large cohort and no ROI restrictions and to compare the ADAD and LOAD signature maps and their ability to detect amyloid burden in cognitively normal groups within and across cohorts. We defined signature maps of cortical thickness separately for LOAD and ADAD by comparing the cognitively impaired and CN groups. The ADAD and LOAD cortical signatures can be accessed online (https://github.com/benzinger-icl/ADcortsig-roi). These cortical signatures included the temporal lobe, parietal, and occipital regions. In the statistical analyses of amyloid, we analyzed only the PC and CN groups. We found the optimal signatures are sensitive in detecting amyloid levels in the cognitively normal groups within their respective cohorts and are significant beyond the inclusion of hippocampus volume, yet the signatures performed poorly when applied cross-cohort (e.g. ADAD signature evaluated in Knight ADRC cohort). In both the Knight ADRC and DIAN cohorts, the signature maps were better predictors when analyzing amyloid as a continuous variable rather than a categorical variable in all statistical models. This result suggests that amyloid levels immediately above and below the amyloid positivity cutoff are important and provide a more sensitive measure to detect the subtle atrophy differences between the CN and PC groups. Finally, by using a continuous variable over dichotomized, our models are optimized for greater statistical power and therefore more able to detect true effects where they exist.

To generate the cortical signatures, we used a vertex-wise approach on a substantial sample size and determined the optimal statistical threshold for multiple comparison correction for each hemisphere. Previous work defining AD cortical signatures used predefined large regions of interest (Dickerson et al., 2009; Jack et al., 2017; Weston et al., 2016) or a similar vertex-wise approach (Wang et al., 2015) with a modest sample size. The spatial extent of significant effects seen in vertex or voxel-wise analyses is highly dependent upon the sample size with a smaller sample creating small peaks which may not translate to other cohorts. Predefined regions of interest from programs like FreeSurfer avoid the small focal effects but incorporate areas that may not be susceptible to AD-related pathology. There is a compromise between having large enough ROIs that the region is stable but not so large it includes marginally predictive areas. By examining cortical maps defined by multiple significance thresholds, we found the optimal tradeoff between these two properties. Each statistical threshold for multiple comparisons correction generated a new cortical map with different ROIs with the strongest threshold (\(p < 0.0001\)) producing the smallest ROIs. We tested how well each statistical threshold for the left and right hemisphere discriminated the PC and CN groups within-cohort and defined optimal cortical signatures from the best performing thresholds. Since our primary interest is focused on individuals transitioning between cognitively normal to early stages of dementia, the selected Monte Carlo thresholds optimize the signatures to the earliest changes in neurodegeneration. We found the optimal selection varied between the left and right hemispheres as well as between the LOAD and ADAD cortical maps. More specifically, the AUROC values in the left hemisphere were overall higher than the right hemisphere for both the ADAD and LOAD cortical maps. Past work have shown hemispherical differences in Alzheimer disease (Kim et al., 2012; Long et al., 2013; Minkova et al., 2017; Toga and Thompson, 2003) and there is also evidence the left hemisphere has an earlier and more rapid progression of atrophy relative to the right hemisphere in AD (Damoiseaux et al., 2009; Thompson et al., 2007). This earlier progression may explain the higher AUROC values in the left hemisphere compared to the right hemisphere. Additionally, the AUROC values in LOAD were higher compared to ADAD, whereas in the regression analyses, the ADAD signature thickness outperformed LOAD. This is primarily due to the inclusion of the age and sex covariates in the regression models. These covariates were not included in the optimization ROC analyses. Since LOAD is largely driven by age, the covariates reduce the predictive value of the cortical signature. This variance shows using a single threshold for each hemisphere and between diseases would have included areas with marginal effect or removed vital areas. Our approach for defining cortical signatures has not, to our knowledge, been attempted previously. We identified the percentage overlap of the cortical signatures on FreeSurfer regions and found the cortical signatures did not encompass any FreeSurfer region entirely, with the highest overlap percentage being 85%. The cortical signature maps are a blend of the most significant areas within each FreeSurfer region, strengthening the point that a larger ROI may not represent the AD cortical thinning behavior. We tested whether different comorbidities, particularly obesity and WMH burden, influence the LOAD cortical signature thickness within the CN\textsubscript{ADRC} and PC\textsubscript{ADRC} groups. BMI had no association to cortical thickness whereas WMH volume had a moderate association, largely due to age. ADAD is also associated with increased WMH burden (Lee et al., 2016). Since the ADAD and LOAD cortical signatures are largely distinct from each other, this may further exemplify WMH burden is not influencing the cortical signatures.

In a comparison between the LOAD and ADAD cortical signatures, we found there is spatial overlap between the two cortical signatures in portions of the right hemisphere inferior and superior parietal regions and portions of the precuneus in both hemispheres (Fig. 3). This spatial overlap may be due to the interrelatedness of these regions on both the structural and functional networks. Further understanding of the underlying network structure in these regions is an important future direction. Examining the significant maps from all possible thresholds (Supplemental Fig. 1, 2) and the effect size maps (Figs. 1B, 2B) show atrophy in ADAD and LOAD is occurring in similar regions, but to a different magnitude. The LOAD signature emphasized atrophy in the temporal lobe regions, while the ADAD signature was most focused on parietal regions. This shift in spatial emphasis is consistent with previous work examining the trophy in LOAD, (La Joie et al., 2012; Singh et al., 2006; Thompson et al., 2003; Whitwell et al., 2007) and ADAD (Apostolova et al., 2011; Cash et al., 2013; Fortea et al., 2010; Gordon et al., 2018A; Knight et al., 2011). The observed atrophy spatial differences also mirror the differences found in the tau pathology patterns, rather than amyloid pathology. Previous work demonstrated tau pathology and cortical atrophy have a high association within the temporal, occipital, parietal, and frontal lobe for the LOAD participants (Gordon et al., 2018A; Knight et al., 2011).
In ADAD, there are higher levels of tau pathology in the prefrontal and a higher precuneus to inferior temporal tau pathology ratio when compared to LOAD (Gordon et al., 2019). Additionally, spatial differences are also seen in early-onset LOAD with previous work corroborating both atrophy (Prisoni, 2005; Möller et al., 2013) and tau PET deposition (Schöll et al., 2017) occur preferentially in the cortex. In contrast, the amyloid spatial pattern between the diseases are largely similar (Shepherd et al., 2009). ADAD having a more prevalent amyloid pathology (Ringenma et al., 2016; Shepherd et al., 2009) as well as an increased amyloid tracer uptake in the basal ganglia (Benzinger et al., 2013; Gordon et al., 2019, 2018a) are the major differences between the diseases.

The exact cause of the different atrophy spatial pattern is unknown but is potentially due to multiple factors. There is evidence that increasing age at dementia onset is tied to more temporal lobe predominate atrophy (Prisoni, 2005; Möller et al., 2013). The pattern may also be tied to the different drivers of pathology in the two forms of the disease. ADAD mutations lead to an overproduction of amyloid burden (Potter et al., 2013) while LOAD is a product of multiple genetic and lifestyle influences that affect both production (Ye et al., 2005) and clearance (Mawuenyega et al., 2010; Moore et al., 2016). Understanding more about this spatial divergence will elucidate unknown factors about AD pathobiology and potentially highlight new therapeutic mechanisms. These different degrees of atrophy between the diseases are most likely playing a role in the unsuccessful cross-cohort analysis in this study and suggests that cortical signature maps are not interchangeable.

This study has many strengths which include the relatively large sample from the Knight ADRC and DIAN cohorts and the application of the cortical signatures to preclinical stages of AD. This comparison between the ADAD and LOAD signatures and the cross-cohort analysis in relation to amyloid pathology has not been studied before to our knowledge. One limitation for this study is the use of cross-sectional data. Identifying regions where longitudinal changes in cortical thickness occur may provide more detail in the progression of the disease. Additionally, future work is needed to examine the performance of the cortical signature in predicting longitudinal cognitive decline. Another limitation is that the cortical signature is only for amnesic presentation due to the use of solely research-based cohorts. Literature suggests that atypical presentations, such as posterior cortical atrophy, might have different atrophy signatures (Lehmann et al., 2011). Clinical cohorts tend to include more participants with comorbidities compared to research cohorts and thus this study may not be translatable to a clinical setting. Finally, prior work suggests there may be differences in atrophy due to specific mutations (Schill et al., 2013), however the infrequency of APP and PSEN2 prevent us from analyzing these differences. As studies of ADAD become more common, it would be of interest to investigate the effect of the different ADAD mutations.

5. Conclusion

The LOAD and ADAD cortical signatures are sensitive to amyloid burden in asymptomatic individuals in their respective cohorts. This finding illustrates that the cortical signature maps are an indicator of AD-related neurodegeneration in early stages of AD and are a useful MRI biomarker measure. However, the different spatial emphasis in atrophy between LOAD and ADAD may be responsible for the unsuccessful cross-cohort analysis. Therefore, the cortical signatures are not compatible across cohorts and should be optimized within the targeted disease.

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CRediT authorship contribution statement

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Declaration of Competing Interests

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: The following authors have no conflicts of interest regarding this study:
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Appendix A. Supplementary data

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