Inferior temporal tau is associated with accelerated prospective cortical thinning in clinically normal older adults

Matthew R. Scott a,b,1, Olivia L. Hampton a,1, Rachel F. Buckley a,b, c, Jasmeer P. Chhatwal a, Bernard J. Hanseeuw e,f, Heidi IL. Jacobs d,e, Michael J. Properzi a, Justin S. Sanchez e, Keith A. Johnson h,e, Reisa A. Sperling a,b, Aaron P. Schultz a,*

a Department of Neurology, Massachusetts General Hospital, Harvard Medical School, Boston, MA, USA
b Center for Alzheimer Research and Treatment, Department of Neurology, Brigham and Women's Hospital, Harvard Medical School, Boston, MA, USA
c Melbourne School of Psychological Science, University of Melbourne, Victoria, Australia
d Faculty of Health, Medicine and Life Sciences, School for Mental Health and Neuroscience, Alzheimer Centre Limburg, Maastricht University, Maastricht, the Netherlands
e Division of Nuclear Imaging and Molecular Imaging, Department of Radiology, Massachusetts General Hospital, Harvard Medical School, Boston, MA, USA
f Faculty of Health, Medicine and Life Sciences, School for Mental Health and Neuroscience, Alzheimer Centre Limburg, Maastricht University, Maastricht, the Netherlands
g Faculty of Health, Medicine and Life Sciences, School for Mental Health and Neuroscience, Alzheimer Centre Limburg, Maastricht University, Maastricht, the Netherlands
h Faculty of Health, Medicine and Life Sciences, School for Mental Health and Neuroscience, Alzheimer Centre Limburg, Maastricht University, Maastricht, the Netherlands
i Faculty of Health, Medicine and Life Sciences, School for Mental Health and Neuroscience, Alzheimer Centre Limburg, Maastricht University, Maastricht, the Netherlands

* Corresponding author. Massachusetts General Hospital/Harvard Medical School, 149 13th Street, Suite 10.036, Charlestown, MA, 02129, United States.
E-mail address: apschultz@mgh.harvard.edu (A.P. Schultz).

1 Both authors contributed equally to this work.

ARTICLE INFO
Keywords: Cortical thinning Tau Longitudinal modeling Flortaucipir PET MRI Alzheimer's disease Imaging

ABSTRACT
Neurofibrillary tau tangles are a hallmark pathology of Alzheimer's disease (AD) and are more closely associated with AD-related cortical atrophy and symptom severity than amyloid-beta (Aβ). However, studies regarding the effect of tau on longitudinal cortical thinning, particularly in healthy aging and preclinical AD, have been limited in number due to the relatively recent introduction of in vivo PET tracers for imaging tau pathology. Here, we investigate [18F]-flortaucipir (FTP, a marker of paired helical filament tau) PET as a predictor of atrophy in healthy aging and preclinical AD. We examine longitudinal structural MRI brain imaging data, retrospectively and prospectively relative to FTP imaging, using piecewise linear mixed-effect models with time centered at each participant's FTP-PET session. Participants include 111 individuals from the Harvard Aging Brain Study who underwent at least three MRI sessions over an average of 4.46 years and one FTP-PET at the approximate midpoint of the observation period. Our primary analyses focus on inferior temporal (IT) FTP standardized uptake value ratios and longitudinal FreeSurfer defined cortical regions of interest. Relationships were also explored using other regional FTP measures (entorhinal, composite, and local), within high and low Pittsburgh compound-B (PiB) PET groups, and with longitudinal subcortical volume. Strong associations between IT FTP and cortical thinning were found, most notably in temporal, midline, and prefrontal regions, with stronger effects generally observed in the prospective as compared to retrospective time frame. Significant differences between prospective and retrospective rates of thinning were found in the inferior and middle temporal gyri, cingulate areas, as well as parietal orbitalis such that higher IT FTP was associated with greater prospective rates of thinning. Within the high PiB group, significant differences between prospective and retrospective rates of thinning were similarly observed. However, no consistent pattern of tau-related change in cortical thickness within the low PiB group was discerned. These results provide support for the hypothesis that tau pathology is a driver of future atrophy as well as provide additional evidence for tau-PET as an effective AD biomarker for interventional clinical trials.

1. Introduction

Longstanding post-mortem studies have identified both neurofibrillary tau tangles and amyloid-beta (Aβ) plaque accumulation as hallmark pathologies of Alzheimer's disease (AD; Braak and Braak, 1991; Braak et al., 2006). Buildup of both pathologies occurs over a long prodromal period (Jack et al., 2013; Villemagne et al., 2013); however, neurofibrillary tangles are more closely associated with atrophy (Mak et al., 2018; Gordon et al., 2018), cognitive decline (Nelson et al., 2012; Maass et al., 2018), and clinical AD phenotype (Jack et al., 2018a). In vivo tau positron emission tomography (PET) studies have shown that tau-PET signal has strong spatial similarities to atrophy patterns (Ossenkoppele

https://doi.org/10.1016/j.neuroimage.2020.116991
Received 18 March 2020; Received in revised form 19 May 2020; Accepted 22 May 2020
Available online 5 June 2020
1053-8119/© 2020 The Author(s). Published by Elsevier Inc. This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/).
Table 1
Demographic information.

|         | Aβ- (%) | Aβ+ (%) | All (%) | Uncorrected p value |
|---------|---------|---------|---------|---------------------|
| N (% subsample) | 72 | 35 | 111 | – |
| Sex(f) | 41 (56.94%) | 19 (54.29%) | 61 (54.95%) | 0.83 |
| APOE4+ | 13 (18.06%) | 22 (62.86%) | 37 (33.33%) | <0.001*** |
| CDR = 0.5 at FTP session | 5 (6.94%) | 1 (2.86%) | 6 (5.41%) | – |
| **Mean (SD)**
| Years of education | 16.28 (3.06) | 16.40 (2.83) | 16.35 (2.98) | 0.84 |
| FTP session MMSE | 29.39 (0.88) | 29.03 (1.04) | 29.25 (0.96) | 0.06 |
| FTP session Logical Memory (DR) | 16.40 (3.97) | 15.11 (3.31) | 16.05 (3.79) | 0.10 |
| Number of MRI follow-ups | 3.55 (0.67) | 3.33 (0.54) | 3.48 (0.63) | 0.10 |
| MRI timespan (years) | 4.36 (1.15) | 4.59 (0.77) | 4.46 (1.05) | 0.32 |
| Retrospective MRI timespan (years) | 2.38 (0.94) | 2.60 (0.81) | 2.43 (0.90) | 0.25 |
| Prospective MRI timespan (years) | 2.10 (0.80) | 1.99 (0.64) | 2.03 (0.78) | 0.51 |
| IT FTP SUVr | 1.43 (0.12) | 1.60 (0.23) | 1.49 (0.19) | <0.001*** |
| EC FTP SUVr | 1.32 (0.22) | 1.64 (0.41) | 1.43 (0.33) | <0.001*** |
| PiB DVR FLR | 1.17 (0.07) | 1.96 (0.42) | 1.43 (0.45) | – |

Participant information is presented at two levels of Aβ burden as well as in the full sample. Aβ group differences were computed using two sample t-tests, chi-squared tests, and were not corrected for multiple comparisons. Four participants did not have Aβ data. FTP and PiB data shown are partial volume corrected. Key: *p < 0.05; **p < 0.01; ***p < 0.001. Abbreviations: Aβ = beta-amyloid; APOE4 = Apolipoprotein E4 genotype; global Clinical Dementia Rating = CDR; Mini-Mental State Examination = MMSE; DR = delayed recall; FTP = flortaucipir; MRI = magnetic resonance imaging; IT = inferior temporal gyrus; EC = entorhinal cortex; SUVr = standardized uptake value ratio; PiB = Pittsburgh compound B; DVR = distribution volume ratio; FLR = frontaltemporal/parietal-retrosplenial.

et al., 2016; Xia et al., 2017; Harrison et al., 2019; Joie et al., 2020) even before clinical AD symptoms arise (Sepulcre et al., 2016). Furthermore, tau-PET signal in these regions has been related to retrospective longitudinal cortical thinning (LaPoint et al., 2017; Gordon et al., 2018; Das et al., 2018), suggesting that atrophy occurs in temporal and spatial proximity to FTP-PET signal. However, the temporal phasing of atrophy versus tauopathy has not yet been fully demonstrated. In this study, we utilize longitudinal structural magnetic resonance imaging (MRI) as well as cross-sectional tau-PET imaging data to investigate retrospective and prospective relationships between in vivo flortaucipir-PET (FTP) signal and cortical thinning in clinically normal (at baseline) older adults. We use inferior temporal (IT) FTP-PET signal as a proxy for AD-related neocortical tau deposition (LaPoint et al., 2017; Sperling et al., 2019; Hanseuwe et al., 2017; Johnson et al., 2016), as the IT gyrus is an early site of tau spread into the neocortex during Braak Stage III (Braak and Braak, 1991; Braak et al., 2006), and autopsy evidence suggests that this spread from medial temporal lobe to neocortex is associated with early cognitive impairment (Braak and Braak, 1997; Hulette et al., 1998; Price and Morris, 1999). In secondary analyses we also explore effects of tau signal in three additional areas (entorhinal cortex [EC], composite, and local FTP), influences of Aβ accumulation, and bearings on subcortical volume.

We first analyze the association between longitudinal thinning over the entire observation period with cross-sectional FTP signal temporally located at the approximate midpoint of the longitudinal MR observation period. We then perform piecewise linear models to separately model retrospective and prospective rates of atrophy while maintaining a common intercept. Examining differences between retrospective and prospective effects, though not explicitly causal, can provide supporting evidence regarding whether FTP signal is more associated with retrospective atrophy, which would be consistent with tau as reflective of atrophy, or prospective atrophy, which would be consistent with tau as a driver of atrophy. This distinction is critical for understanding how to use and interpret FTP-PET as a biomarker for AD and whether tau is a good candidate target for intervention.

2. Methods
2.1. Participants

Participants included 111 individuals from the Harvard Aging Brain Study (HABS), a longitudinal study of normal aging and preclinical AD conducted at Massachusetts General Hospital, Athinoula A. Martinos Center for Biomedical Imaging (Dagley et al., 2017). We refer to pre-clinical AD in the sense of a cognitively normal, elderly cohort that can be characterized in terms of NIA-AA criteria (Jack et al., 2018b; Sperling et al., 2011). All participants provided informed consent and were studied under protocols approved by the Partners Human Research Committee. Participants were screened for unstable medical or neuropsychological diseases. At study entry, all participants were assessed as clinically normal (global Clinical Dementia Rating [CDR] = 0, MMSE > 25, and performance within 1.5 standard deviations of age and education-adjusted norms on the Logical Memory Delayed Recall), without current clinical depression (Geriatric Depression Scale < 11), or active psychiatric illness (Folstein et al., 1975; Morris, 1993; Wechsler, 1987; Yesavage et al., 1982). Furthermore, participants were also excluded from the study if they had a history of alcoholism, drug abuse, or head trauma. Diagnosis was evaluated in consensus meetings using longitudinal clinical and neuropsychological assessments without regard for AD imaging biomarkers. All participants were clinically normal at the time of the FTP-PET scan.

Sample characteristics are defined in Table 1. 11 participants progressed to a global CDR of 0.5 at their latest MRI session. Neuropsychological evaluation approximately 1.5 years after each participant’s final MRI also showed a total of 20 participants progressing to a global CDR of 0.5 and 1 participant progressing to 1.0. There was an overlap of 73 participants in this sample compared to previous work on retrospective cortical thickness from HABS (LaPoint et al., 2017). The inclusion criteria for retrospective MRIs in the work of LaPoint included MRI sessions which were collected within one year following the tau-PET scan (in addition to scans before). As such, the current work included more prospective MRI sessions and classified timepoints temporally in relation to the tau-PET scan instead of labelling nearest MRI scans to the tau scan as contemporaneous. Lastly, all individuals in the current sample were required to have at least three MR sessions, which included both a retrospective and a prospective MR session relative to the tau-PET session.

2.2. Longitudinal MRI

MRI was performed on a TimTrio 3.0T imaging system (Siemens Medical Systems, Erlangen, Germany) with a 12-channel phased-array head coil over an average range of 4.46 years. Head motion was controlled with a foam pillow and extendable padded head clamps. Scanner noise was attenuated with ear plugs. Structural T1-weighted anatomical images were acquired using a magnetization-prepared rapid
acquisition gradient-echo (MPRAGE; ADNI GO/2 accelerated) sequence with the following parameters: repetition time (TR) = 2300 ms, echo time (TE) = 2.95 ms, inversion time = 900 ms, flip angle = 9°, 1.05 × 1.05 × 1.2 mm resolution. However, 28 of the 386 sessions had a different sequence (ADNI GO/2 unaccelerated) with the following parameters: TR = 2300 ms, TE = 2.98 ms, inversion time = 900 ms, flip angle = 9°, 1.00 × 1.00 × 1.2 mm resolution. These sequences have been deemed equivalent. T1-weighted images were then processed with FreeSurfer version 6.0 to identify gray-white as well as pial surfaces and to produce an automatic region of interest (ROI) parcellation (Desikan et al., 2006; Fischl, 2012).

Following previously described cross-sectional quality control measures (Dagley et al., 2017), each participant's images were run through the FreeSurfer longitudinal processing stream (Reuter et al., 2012). In the longitudinal processing stream, a temporally unbiased template is created for each participant based on all MR timepoints. Next, each time point is resampled to a median template space, with the intention of reducing random variability between participant timepoints and improving the sensitivity of analyses. Identical manual quality control of FreeSurfer reclones processed through the longitudinal pipeline was repeated to ensure accuracy of data, including skull stripping, white matter edits, and placement of control points in the templates and longitudinally processed scans as needed. All 34 cortical thickness ROIs produced by FreeSurfer were utilized in our analyses (Desikan-Killiany). Only bihemispheric averages for each region were included in analyses as we had no a priori hypotheses on laterality. Effects on longitudinal lateral ventricle, inferior lateral ventricle, thalamus proper, caudate, putamen, pallidum, brain stem, hippocampus, amygdala, accumbens, cerebrospinal fluid, total gray, cortex, and ventral diencephalon volume (mm³; based on FreeSurfer subcortical segmentation) are reported in addition to cortical ROIs.

2.3. Tau-PET imaging

[18F]-FTP was prepared with a radiochemical yield of 14 ± 3% and specifically activity of 216 ± 60GBq/μmol at the end of synthesis as previously reported (Johnson et al., 2016), and validated for human use (Shoup et al., 2013). Data were acquired using a Siemens/CTI ECAT HR + scanner (3-dimensional mode, 63 image planes, 15.2 cm axial field of view, 5.6 mm trans axial resolution, and 2.4 mm slice interval), with acquisition taking place 80–100 min for 75 sessions and 75–105 min for 36 sessions, after a 9.0–11.0 mCi bolus injection in 4 × 5 min frames for 80–100 min sessions and 6 × 5 for 75–105 min sessions. FTP-PET data were reconstructed as well as attenuation corrected, and each frame was evaluated to verify adequate count statistics as well as absence of head motion.

To evaluate the anatomy of cortical FTP binding, PET scans were rigidly coregistered to the individual’s MPRAGE images using SPM12 (Function Imaging Laboratory, Wellcome Department of Cognitive Neurology, London, UK). ROIs defined by the cross-sectional FreeSurfer analysis at the MRI session closest to the FTP scan were transformed into the PET native space. Reported FTP data were partial volume corrected (PVC) using the Geometric Transfer Matrix (GTM; Rouset et al., 1998) method implemented in FreeSurfer version 6.0 (Greve et al., 2016). Associations involving FTP were validated with non-PVC data. FTP signal was expressed in FreeSurfer ROIs as standardized uptake value ratios (SUVr), using cerebellar gray as the reference. Effects of a composite ROI (consisting of amygdala, fusiform, IT, middle temporal, and EC), EC, and local (i.e., both FTP and thickness measures were obtained from the same ROI) FTP are reported as supplemental information.

2.4. Amyloid-PET imaging

Aβ-PET imaging data were acquired on average 79 days from each participant's FTP-PET visit (SD = 77.80 days; median = 54 days). Deposition of Aβ was measured with PET using Pittsburgh compound-B (11C-PiB) according to previously described methods (Johnson et al., 2007). After injection of 8.5–15.0 mCi PiB, we acquired 60 min of dynamic PET data using the Siemens/CTI ECAT HR+ PET scanner (see Tau-PET Imaging for parameters; 39 frames: 8 × 15s, 4 × 60s, and 27 × 120s. PiB-PET data were reconstructed as well as attenuation corrected, and each frame was evaluated to confirm adequate count statistics as well as absence of head motion. Using Logan graphical analysis, PiB retention was expressed as a distribution volume ratio (DVR) using a cerebellar gray reference region (Price et al., 2005). Neocortical Aβ deposition was quantified using an aggregate DVR from a set of regions that comprised most of the association cortex, including frontal, lateral parietal and temporal, as well as retrosplenial cortices (FLR). Reported PiB data were PVC using the GTM method; however, PiB was only used in a secondary analysis to subset the sample into high and low Aβ subgroups. A FLR threshold of 1.34 was derived from a Gaussian mixture model on the full HABS cohort, as described by Mormino et al. (2014a). Dichotomous PVC and non-PVC classification differed by only two participants.

2.5. Statistical analyses

Statistical analyses were performed in MATLAB R2018a (MathWorks, Inc., Natick, MA) using linear mixed-effects models with random intercepts for each participant and random time slopes. Regional bihemispheric thickness was the dependent variable in all analyses (lateral patterns are reported as supplemental material). As the purpose of the analysis is to understand atrophy with respect to FTP-PET signal, time was centered at each participant’s FTP-PET visit (i.e., the time of each participant’s FTP-PET session is 0). Time between the PET scan and the closest MRI scan did not significantly alter the results of our analyses. Additionally, all models included age, sex, and their interactions with time as covariates. We investigated the effect of FTP signal on cortical thickness over all timepoints (Table 2, Model 1). Thereafter, the effect of FTP on retrospective and prospective thinning was analyzed using a piecewise linear regression approach (Table 2, Model 2B). Significant differences between retrospective and prospective slopes were assessed via the interaction term of FTP × time × observation period (i.e., retrospective vs prospective relative to the FTP scan; Table 2, Model 2A). This approach allows for both naive modeling of non-linearities and comparisons between retrospective and prospective time periods (relative to the FTP-PET measurement). All results are reported at a threshold of p < 0.05, uncorrected, to investigate the pattern of these relationships across the cortex, then False Discovery Rate (FDR) corrected to confirm which associations survived adjustment for multiple comparisons. Estimates reported are not standardized. To exemplify findings, model fits of
IT FTP effects on prospective and retrospective IT, EC, and cuneus thinning were generated using estimates from Model 2. Analyses were also repeated in both low and high PiB groups separately. This was done to exemplify the findings separately in healthy aging and preclinical AD cohorts as well as to avoid statistical complications and interpretability issues when modeling interactions between collinear PiB and FTP measures.

Our hypothesis is directional, such that cortical thickness is expected to decrease over time; however, as previous accounts of neuroinflammatory processes have been reported, the possibility that increases in thickness occur is not ruled out (Heneka et al., 2015; Fortea et al., 2010; Sala-Llonch et al., 2015; Weston et al., 2016). Effects across all ROIs are reported regardless of directionality. In addition to primary IT FTP analyses, we conducted additional analyses: 1) Effects of EC FTP, composite FTP, and local FTP on cortical thinning were investigated in the same manner as IT FTP. 2) We explored effects of IT FTP signal separately in high and low PiB groups to better reflect aging versus preclinical AD. 3) The associations of IT FTP signal and subcortical volume were characterized across 14 regions. Hippocampal volume is highlighted as it is a well characterized AD biomarker of brain atrophy.

3. Results

3.1. Inferior temporal tau

The effect of IT FTP on longitudinal thinning across the full observation period (Model 1, IT FTP X Time; Fig. 1, Thickness Slope) was significant in sixteen regions before FDR correction and eleven regions after. Effects were observed in a variety of brain areas, however, six of the nine temporal ROIs explored had the largest effects of all 34 ROIs (in order of increasing t-statistic magnitude): EC, temporal pole, parahippocampal, fusiform, IT, and middle temporal.

For the retrospective observation period (Model 2B, IT FTP:Time;G1; Fig. 1, Retrospective Thickness Slope), higher IT FTP signal was related to faster rates of retrospective thinning in paracentral (t(376) = -2.14, p = 0.037, p < 0.05) and EC regions (t(376) = -2.58, b = 0.05, p < 0.05). Caudal anterior cingulate (t(376) = 2.45, b = 0.03, p < 0.05) conversely exhibited a significant increase in retrospective thickness rates with higher IT FTP signal. However, no regions survived correction for multiple comparisons. For the prospective observation period (Model 2B, IT FTP:Time:G2; Fig. 1, Prospective Thickness Slope), higher IT FTP signal was significantly associated with faster rates of cortical thinning in eighteen regions before correction for multiple comparisons and thirteen regions following. Temporal regions were prominent, with IT (t(376) = -4.21, b = -0.05, p < 0.001) and middle temporal gyrus (t(376) = -4.34, b = -0.06, p < 0.001) exhibiting the largest t-statistics. Cingulate regions such as the rostral anterior cingulate (t(376) = -2.77, b = -0.04, p < 0.01) and posterior cingulate (t(376) = -3.07, b = -0.03, p < 0.01) also showed potent effects, along with the pars orbitalis (t(376) = -3.63, b = -0.06, p < 0.001).

Differences between retrospective and prospective rates of thinning (Model 2A, IT FTP X Time X G; Fig. 1, Slope Differential) were significant in IT (t(376) = -2.04, b = -0.04, p < 0.05), middle temporal (t(376) = -2.11, b = -0.04, p < 0.05), posterior cingulate (t(376) = -2.19, b = -0.03, p < 0.05), rostral anterior cingulate (t(376) = -2.53, b = -0.06, p < 0.05), and pars orbitalis (t(376) = -2.73, b = -0.07, p < 0.01) such that higher IT FTP signal was associated with faster prospective thinning compared to retrospective thinning. Caudal anterior cingulate (t(376) = -3.20, b = -0.07, p < 0.01) showed significantly stronger prospective thinning by IT FTP signal, driven by both significant increases in retrospective thickness and significant prospective thinning. No ROIs survived FDR correction.

T-statistics of the above analyses are displayed on brain maps in Fig. 2. These analyses were performed with non-PVC FTP data and no differences in significant ROIs (compared to PVC FTP) were observed following FDR correction. Local, EC, and composite FTP signals were analyzed in the same manner as IT FTP, and effects on cortical thinning are reported in Supplemental Figs. 1 and 2. Lateralized thickness patterns are reported in Supplemental Fig. 3. Spaghetti plots of thickness in IT, EC, and composite FTP signals are displayed on brain maps in Fig. 2. These analyses were performed with non-PVC FTP data and no differences in significant ROIs (compared to PVC FTP) were observed following FDR correction. Local, EC, and composite FTP signals were analyzed in the same manner as IT FTP, and effects on cortical thinning are reported in Supplemental Figs. 1 and 2. Lateralized thickness patterns are reported in Supplemental Fig. 3. Spaghetti plots of thickness in IT, EC,
caudal anterior cingulate, and cuneus across the full follow-up range and grouped by median IT FTP signal are shown in Fig. 3.

3.2. Amyloid

We repeated the above analyses separately in participants with low Aβ (putatively normal aging) and high Aβ (preclinical AD), stratified by an aggregate DVR of PiB-PET signal. Effects within each group are reported in Fig. 4. The low PiB group exhibited lower IT FTP signal than the high PiB group (t(105) = 4.89, p < 0.001; Table 1). In the low PiB group, significant associations between IT FTP signal and cortical thickness slopes across the full, retrospective, or prospective were only observed in the direction of increased thickness (Fig. 4A). Although we observed a significant difference in slopes in the low PiB group within IT (t(245) = -
1.97, b = 0.07, p < 0.05), inferior parietal (t(245) = -2.00, b = -0.05, p < 0.05), parahippocampal (t(245) = -2.03, b = -0.05, p < 0.05), and pericalcarine (t(245) = 2.04, b = 0.09, p < 0.05) regions, these effects were accentuated by positive retrospective slopes that were not significantly different from 0 (except for pericalcarine). No ROIs in any low PiB analyses survived FDR correction. The high PiB group exhibited significant associations between IT FTP signal and rates of cortical thinning (Fig. 4B) in full, retrospective, or prospective periods. Retrospective increases in thickness were observed in medial orbitofrontal (t(108) = 2.02, b = 0.03, p < 0.05) and pars orbitalis (t(108) = 2.16, b = 0.04, p < 0.05). Rates of thinning were generally higher prospectively rather than retrospectively, with six regions demonstrating a significant difference between retrospective and prospective thinning estimates. Of these six, only the pars orbitalis survived FDR correction (t(108) = -4.49, b = -0.15, p < 0.001).

3.3. Subcortical volume

We then investigated the association between longitudinal hippocampal volume and IT FTP signal. We observed significant effects of IT
FTP signal across the full follow-up ($t(378) = -2.67$, $b = -39.44$, $p < 0.01$) and prospective period ($t(376) = -3.22$, $b = -72.88$, $p < 0.01$), but no effect was found retrospectively. Differences between retrospective and prospective hippocampal volume rates were marginal ($t(376) = -1.83$, $b = -55.13$, $p = 0.06$). When separately analyzing low and high PiB groups, slope differentials were found in low PiB participants ($t(245) = -2.04$, $b = -112.56$, $p < 0.05$) but not high PiB participants ($t(108) = 0.05$, $b = 2.03$, $p = 0.96$). No significant effects were observed in either group over the full, retrospective, or prospective observation period.

Effects of IT FTP signal across subcortical ROIs are shown on brain renderings in Fig. 5. Not presented in Fig. 5, greater IT FTP signal was significantly associated with total gray ($t(378) = -2.52$, $b = -3.15 \times 10^3$, $p < 0.05$) and cortex ($t(378) = -2.77$, $b = -2.97 \times 10^3$, $p < 0.01$) atrophy over the full follow-up range, and putamen volume was marginal ($0.05 < p < 0.10$). Higher IT FTP signal was found to be significantly associated with prospective subcortical atrophy across the cortex ($t(376) = -3.04$, $b = -5.99 \times 10^3$, $p < 0.01$) and total gray ($t(376) = -3.29$, $b = -7.14 \times 10^3$, $p < 0.01$). At larger IT FTP levels, prospective rates of atrophy were faster than retrospective rates with total gray volume ($t(376) = -2.02$, $b = -6.22 \times 10^3$, $p < 0.05$). Brain stem, hippocampus, and thalamus exhibited marginal slope differentials ($0.05 < p < 0.10$).

4. Discussion

The present study characterized the associations of FTP-PET signal in the IT gyrus with both retrospective and prospective longitudinal cortical thinning across 34 ROIs, as well as subcortical atrophy in 14 ROIs, in a clinically normal (at baseline) sample from the Harvard Aging Brain Study. We observed that higher IT FTP signal is associated with accelerated rates of cortical thinning, and that these effects are generally stronger in prospective than retrospective observation periods. These findings are consistent with prior reports and the hypothesis that tau pathology is a driver of atrophy rather than a reflection of atrophy (Sabuncu and Konukoglu, 2014; Thompson et al., 2003; LaPoint et al., 2017; Gordon et al., 2018; Mak et al., 2018; Harrison et al., 2019).

There were widespread effects of IT FTP signal on longitudinal cortical thinning, but temporal regions were the most notable of these by surviving correction for age, sex, and multiple comparisons. Retrospective thinning was found in paracentral and EC regions. A marginal effect of the parahippocampus ($p = 0.05$) was observed, which was significant in both left and right hemispheres in the previous work from HABS (LaPoint et al., 2017). Any differences in effect sizes between the current work and those found by LaPoint can be attributed to the current work’s overlapping but different set of participants, characterization of retrospective MRIs (see Methods section), utilization of bihemispheric ROIs, and an updated FreeSurfer pipeline.

Prospective associations were more prevalent, with over half the ROIs exhibiting a significant prospective effect, particularly in cingulate and temporal areas. Significant differences between retrospective and prospective slopes were found in the posterior cingulate, IT gyrus, middle temporal gyrus, rostral anterior cingulate, and pars orbitalis, such that higher levels of IT FTP signal were associated with faster thinning prospectively as compared to retrospectively. These results align with prior reports highlighting temporal and medial regions as early sites for atrophy along the AD trajectory (Braak and Braak, 1991; Chan et al., 2001; Killiany et al., 2002; Scallill et al., 2002; Dickerson et al., 2009; Mormino et al., 2014b; Mak et al., 2018; Das et al., 2018; Harrison et al., 2019). Differential slopes were also found in inferior lateral and lateral ventricles, providing further support to our cortical findings as significant volumetric increases in the ventricles could correspond to decreases in thickness or volume of neighboring areas. Moreover, we observed a $p = 0.06$ when using hippocampal volume. Caudal anterior cingulate thickness was also found to exhibit differential rates, and its relationship was driven by a significant increase in retrospective thickness. Fig. 6 illustrates these relationships at varying levels of IT FTP in characteristic regions.

As expected, the spatial effects of EC FTP signal on longitudinal thinning were similar to that of IT FTP, which is not surprising given the high degree of collinearity between EC and IT FTP signal ($r = 0.83$, $p < 0.001$), and given that EC is a location of both age and AD related tauopathy (Price, 1993). Both IT and EC FTP had significant retrospective effects on EC thinning, and their prospective effects differed only slightly. Of note, we observed a strong and consistent relationship between IT FTP and EC thinning in both retrospective and prospective periods (the same pattern is true for EC FTP), which may suggest that we have missed the tipping point of the pathological cascade for the EC in the HABS sample. However, when looking at the effect of IT FTP on other medial and lateral temporal regions, we see a marked increase in the association between IT FTP signal in the prospective time frame, relative to the retrospective time frame. This may indicate that lateral temporal regions represent an inflection point in preclinical AD, and that early increases in FTP-PET signal in the IT gyrus are associated with cortical thinning in a broader AD type pattern. Analyses of local FTP signal provide additional evidence for this hypothesis, as IT thinning was the only region that exhibited a significant slope difference.

Studies of preclinical familial AD have reported an unexpected initial increase in cortical thickness, prior to later decline, indicating a potential preclinical inflammatory process (Fortea et al., 2010; Sala-Llonch et al., 2015). Though our hypothesis was directional in that we predicted increases in IT FTP to be associated with longitudinal thinning, the caudal anterior cingulate exhibited effects of IT FTP in the direction of increasing retrospective thickness. Increases in thickness were also observed in both high and low PiB groups, though low PiB effects may not be AD-related and may reflect partial volume effects. As none of these effects survived FDR correction, more work investigating early neuronal inflammatory processes is needed.

We ran our IT FTP models separately in individuals with high or low PiB-FTP signal. In the low PiB group, we did not observe significant thinning over the full, retrospective, or prospective observation period, as only increases in thickness were observed. As expected, high PiB participants exhibited thinning effects over the full observation period, retrospectively, and prospectively. Differences between retrospective and prospective thickness were seen in both low and high PiB groups, yet these effects were largely driven by positive slopes in the retrospective time frame. Within the high PiB group, significant effects over the full and prospective period as well as significant rate differences largely mirrored the findings from the whole sample. The only regions in the high PiB individuals which varied from the full sample in differences in slopes were the IT and frontal pole. These relationships indicate that CN participants with elevated levels of both tau and amyloid pathologies represent a heightened risk group, a finding that has been highlighted in recent prospective studies investigating cognitive trajectories within this same cohort (Sperling et al., 2019), as well as other studies investigating the association of amyloid and tau pathology to longitudinal thinning (Gordon et al., 2018; Mak et al., 2018). It should also be noted that FTP-PET and PiB-FTP signals are correlated, and that high levels of IT FTP signal are rare in low PiB individuals.

There are additional limitations of the current study to consider. First, our FTP-PET measures are cross-sectional, meaning we do not know when FTP-PET signal began increasing or at what rate it is currently increasing, which means that our results are not direct evidence of a particular temporal ordering relative to atrophy and no causal inferences may be drawn directly from these results. Additional follow-up work utilizing longitudinal FTP-PET and longitudinal structural imaging is needed in both concurrent and non-overlapping time frames. Regarding effects in high amyloid versus low amyloid individuals, we must acknowledge the collinearity between PiB and FTP measurements and the general lack of IT FTP-PET signal in the low amyloid group. Additionally, as our primary analysis examined three-way interactions of time, IT FTP, and temporal grouping (Model 2), we chose not to explore four-way interactions with continuous PiB to avoid interpretation.
Fig. 5. Subcortical Volume

3D brain renderings of significant (no FDR correction) IT SUVr t-statistics. In order of increasing magnitude (A) Subcortical volume over the full follow-up range in the hippocampus, lateral ventricles, amygdala, and inferior lateral ventricles. (B) Prospective subcortical volume in the pallidum, putamen, thalamus proper, amygdala, hippocampus, lateral ventricles, and inferior lateral ventricles. (C) Retrospective subcortical volume in the brain stem. (D) Slope differences between retrospective and prospective thickness in the lateral and inferior lateral ventricles. Abbreviations: IT = inferior temporal gyrus; SUVr = standardized uptake value ratio; FTP = flor- taucipir; FDR = False Discovery Rate.
complexity. Thus, we opted to use a dichotomized PiB measure to provide more straightforward interpretability of the results. Future work could explore interactive effects of continuous PiB, as well as potential non-linearities involving PiB.

Our findings show that significant differences between retrospective and prospective rates of cortical thinning are associated with IT FTP-PET signal in AD vulnerable regions, suggesting that elevated IT FTP may be prognostic of accelerating rates of atrophy. Additional studies are needed to examine how these relationships change with longitudinal PET data, cognition, and disease progression. Our findings are consistent with the hypothesis that tau is a driving agent of cortical thinning and suggest that FTP-PET holds promise as both a biomarker for AD progression and a monitor of therapeutic effects in interventional trials.

Declaration of competing interest

Dr. Schultz has received grants from the National Institute on Aging and served on paid advisory boards for Janssen Pharmaceuticals (2013) and Biogen (2016). Dr. Hanseeuw is funded by the Belgian National Fund for Scientific Research (FNRS) #SPD28094292. Dr. Jacobs reported funding from the European Union’s Horizon 2020 Research and Innovation Programme under the Marie Skłodowska-Curie Grant agreement (IF-2015-GF, 706714). Dr. Johnson has served as a paid consultant for Bayer, GE Healthcare, Janssen, Siemens Medical Solutions, Genzyme, Novartis, Biogen, Roche, AZTherapy, GEHC, Lundbeck, Genentech, Lilly/Avid, AC Immune, and Abbvie. Dr. Sperling has received grants from National Institute on Aging, Alzheimer's Association, Eli Lilly and Co., and Janssen Pharmaceuticals. She has consulted for AC Immune, Eisai, National Institute on Aging, Alzheimer’s Association, Eli Lilly and Co., Avid, AC Immune, and Abbvie. Dr. Sperling has received grants from the National Institute on Aging, Alzheimer’s Association, Eli Lilly and Co., Avid, AC Immune, and Abbvie. Dr. Sperling has received grants from the National Institute on Aging, Alzheimer’s Association, Eli Lilly and Co., Avid, AC Immune, and Abbvie. Dr. Sperling has received grants from the National Institute on Aging, Alzheimer’s Association, Eli Lilly and Co., Avid, AC Immune, and Abbvie. Dr. Sperling has received grants from the National Institute on Aging, Alzheimer’s Association, Eli Lilly and Co., Avid, AC Immune, and Abbvie. Dr. Sperling has received grants from the National Institute on Aging, Alzheimer’s Association, Eli Lilly and Co., Avid, AC Immune, and Abbvie.

Acknowledgements

We would like to acknowledge the participants of the Harvard Aging Brain Study for their dedication. This work was supported with funding from National Institutes of Health grants via R01AG027435 and for the Harvard Aging Brain Study (P01AG036694) and R01AG046396 (Dr. Johnson). This research was carried out at the Athinoula A. Martinos Center for Biomedical Imaging at the Massachusetts General Hospital, using resources provided by the Center for Functional Neuroimaging (P41EB015896), a P41 Biotechnology Resource Grant supported by the National Institute of Biomedical Imaging and Bioengineering (NIBIB), National Institutes of Health. This work also involved the use of instrumentation supported by the NIH Shared Instrumentation Grant Program and/or High-End Instrumentation Grant Program; specifically, S10RR021110, S10RR023401, S10RR019307, S10RR019254, and S10RR023043.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.neuroimage.2020.116991.
