Deconstructing pathological tau by biological process in early stages of Alzheimer disease: a method for quantifying tau spatial spread in neuroimaging

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Summary

Background Neuroimaging studies often quantify tau burden in standardized brain regions to assess Alzheimer disease (AD) progression. However, this method ignores another key biological process in which tau spreads to additional brain regions. We have developed a metric for calculating the extent tau pathology has spread throughout the brain and evaluate the relationship between this metric and tau burden across early stages of AD.

Methods 445 cross-sectional participants (aged ≥ 50) who had MRI, amyloid PET, tau PET, and clinical testing were separated into disease-stage groups based on amyloid positivity and cognitive status (older cognitively normal control, preclinical AD, and symptomatic AD). Tau burden and tau spatial spread were calculated for all participants.

Findings We found both tau metrics significantly elevated across increasing disease stages (p < 0.0001) and as a function of increasing amyloid burden for participants with preclinical (p < 0.0001, p = 0.0056) and symptomatic (p = 0.010, p = 0.0021) AD. An interaction was found between tau burden and tau spatial spread when predicting amyloid burden (p = 0.00013). Analyses of slope between tau metrics demonstrated more spread than burden in preclinical AD (β = 0.59), but then tau burden elevated relative to spread (β = 0.42) once participants had symptomatic AD, when the tau metrics became highly correlated (R = 0.83).

Interpretation Tau burden and tau spatial spread are both strong biomarkers for early AD but provide unique information, particularly at the preclinical stage. Tau spatial spread may demonstrate earlier changes than tau burden which could have broad impact in clinical trial design.

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Keywords: Alzheimer disease; Positron emission tomography; Tau spread; Tau propagation

Introduction Alzheimer disease (AD) is a neurodegenerative disease characterized by the accumulation of amyloid-beta plaques and tau neurofibrillary tangles (NFTs) as well as subsequent neurodegeneration and cognitive decline.1-4 Amyloid-beta has long been thought to be a driving initial factor of AD5 as plaques begin developing,5,3 decades before the presentation of clinical symptoms.6-8 The presence of NFTs occurs later in the disease course and is shown to be tightly coupled with neurodegeneration and cognitive impairment.11-13 Given its prominent role in the transition to an impaired state understanding tauopathy is of utmost importance.

Early histopathological studies found a progressive spatial pattern for NFTs beginning in the entorhinal...
cortex, then spreading to the temporal cortex and finally isocortical regions following Braak-staging. This pattern of spread occurs after amyloid deposition and aligns with widely-accepted cognitive impairments in AD, typically beginning with memory deficits and progressing to executive, visuospatial, and language dysfunction. Tau PET studies have provided in vivo confirmation of neuropathology findings showing distinct spatiotemporal correlations of tau to regional cortical atrophy and resulting cognitive dysfunction. Tau PET uniquely allows for the in vivo assessment of spatiotemporal progression, allowing us to evaluate the simultaneous spread of tau pathology to new brain regions and increasing tau levels in previously affected regions. The progression of tau can be characterized by both increasing accumulation of NFTs (tau intensity) and the transportation of tau throughout the brain (tau spatial spread).

Spatial patterns of tau may be explained by prion-like tau seeding of misfolded proteins along synaptically-connected neuronal circuits. Transneuronal tau propagation has been supported in cellular research, mouse models, post-mortem studies, and white matter connectivity. This process of spread is activity-dependent, with increased propagation at higher neuronal firing rates, resulting in strong correlations of tau within functional networks.

Neuroimaging AD research typically measures global tau burden either by defining a threshold for tau or by averaging tau PET signal across a pre-defined set of regions of interest (ROIs). These summary measures, typically focused on the temporal lobe, capture areas of early sequential tau deposition, but aggregate only a small set of regions rather than specifically considering the degree of spatial spread through the cortex. Some studies have tried to account for spread by evaluating tau PET signal separately within several ROIs while others have been able to model the progression of tau pathology to new regions. These analyses do not, however, distinctly differentiate between nor distinctly compare tau intensity and tau spatial spread throughout AD. They additionally force a specific regional structure which is ill suited to subject variability and atypical presentations. Some studies have attempted to address the issue of inter-individual differences and atypical tau spatial patterns but continue to focus on identifying new ROIs and summary measures rather than evaluating tau spread as a separate metric.

Disentangling tau intensity and tau spatial spread is important in elucidating which of these biological processes is the driving factor for cognitive decline in AD or whether the amount of tau pathology and spread to other regions interact throughout the disease. With many potential tau-targeting drugs currently being tested in clinical trials, evaluating these components of tau concurrently could be critical for evaluating trial success as well as deciding which drug to deliver. This paper proposes a method for quantifying global tau spatial spread extent, independent from the specific underlying mechanism of protein movement, and characterizes the relationship between tau intensity and tau spatial spread using tau PET in the early stages of AD to determine the efficacy of each biological process as a biomarker for early AD progression.

**Methods**

**Participants**
Participants enrolled in ongoing studies of memory and aging from the Charles F. and Joanne Knight Alzheimer Disease Research Center (Knight ADRC) at Washington University School of Medicine (WUSM) between 2014 and 2020 were used in our cross-sectional analyses. Inclusion criteria included age 50 and older with amyloid and tau...
PET, structural magnetic resonance imaging (MRI), and Clinical Dementia Rating® (CDR®). All data were collected within a one-year period per participant.

Similar data from a cohort of younger controls (YC, age ≤ 49) were chosen from both the Knight ADRC and Dominantly Inherited Alzheimer Network (DIAN)-Observational study at WUSM.49 All YC from the Knight ADRC were amyloid-negative and cognitively normal as determined by corresponding amyloid PET and CDR. YC from DIAN were amyloid-negative and cognitively normal as well as non-carriers for the PSEN1, PSEN2, and APP genetic mutations studied in DIAN.

Final sample size included all participants from the Knight ADRC and DIAN cohorts who met inclusion criteria, resulting in 445 older participants and 21 younger controls. Participant demographic information was self-reported.

Ethics
All participants provided written informed consent and the process for data collection was approved by the Washington University Human Research Protection Office, which serves as the central institutional review board (IRB), for the Knight ADRC and DIAN studies (protocols 201106339, 201306009, 201106168, 201106148, and 201409014).

Imaging acquisition and processing
T1-weighted MRI scans were acquired on a DIAN-approved 3T scanner at a resolution of either $1 \times 1 \times 1.25$ mm$^3$ or $1 \times 1 \times 1$ mm$^3$. Cortical and subcortical ROIs for PET analyses were defined from the structural T1 using FreeSurfer (v5.3-HCP: http://surfer.nmr.mgh.harvard.edu/).62

Amyloid PET imaging was performed using 10.01 ± 0.61 mCi of $^{18}$F-florbetapir ($^{18}$F-AV-45) or 14.94 ± 3.85 mCi of $^{11}$C-Pittsburgh Compound B ($^{11}$C-PiB). Tau PET imaging was performed using 9.04 ± 0.86 mCi of $^{18}$F-flortaucipir ($^{18}$F-AV-1451). For $^{18}$F-AV-45 and $^{18}$F-AV-1451, regional and voxel-specific standard uptake value ratios (SUVRs) were calculated using the cerebellar grey as the reference region for the 50–70 minute and 80–100 minute post-injection window, respectively, using the FreeSurfer-based PET Unified Pipeline (PUP; https://github.com/ysu001/PUP).63 For $^{11}$C-PiB, the post-injection window for SUVR quantification was dependent on the cohort source (30–60 minute for Knight ADRC and 40–70 minute for DIAN).

Global amyloid burden was evaluated with a cortical summary measure by averaging partial volume corrected SUVR across precuneus, prefrontal cortex, gyrus rectus, and lateral temporal cortex ROIs.62 Amyloid positivity was determined using our previously published thresholds for $^{18}$F-AV-45 (SUVR > 1.19) and $^{11}$C-PiB (SUVR > 1.42).62,64

Centiloid is a commonly used method in the field for standardizing amyloid quantification across tracers and research centers.65 The cortical summary measures for $^{18}$F-AV-45 and $^{11}$C-PiB were converted to Centiloids per our published equations.64

Clinical testing
Cognitive status was evaluated using the CDR, a clinical tool for assessing the presence and, when present, the severity of Alzheimer dementia. The CDR is calculated based on scores as to whether there has been a change from previously attained levels of function in 6 domains—memory, orientation, judgment and problem solving, community affairs, home and hobbies, and personal care. The global CDR is scored on a scale 0–3 that denotes no impairment (CDR = 0), very mild impairment (CDR = 0.5), mild impairment (CDR = 1), moderate impairment (CDR = 2), and severe impairment (CDR = 3).

Disease-stage group classification
Participants were assigned to disease-stage groups dependent on amyloid positivity (A+) and cognitive symptoms (CDR), resulting in three final groups for analysis: Aβ-CDR0 (older control, OC), Aβ+CDR0 (preclinical AD), and Aβ+CDR+0 (symptomatic AD). Disease-stage groups correspond to early clinical progression of AD, spanning the continuum of amyloid-beta plaque accumulation and cognitive decline.62

Tau Index (TI) and Tau Spatial Spread (TSS)
Tau index (TI) was calculated with PUP-processed tau PET ROI outputs as the mean regional SUVR for the four regions previously identified to characterize early tau accumulation (entorhinal cortex, amygdala, lateral occipital cortex, and inferior temporal cortex).66 TI was calculated for participants as a measure of tau intensity.

Tau spatial spread (TSS) was measured from voxel-wise tau PET images as the proportion of the relevant brain voxels with abnormal tau pathology (Fig. 1). Image processing and computations specific to the calculation of TSS were conducted with PUP-processed participant files and FMRI B Software Library (FSL v6.0; https://fsl.fmrib.ox.ac.uk/).

Participant scans were first preprocessed and co-registered for voxel-wise analyses. T1 MR image, tau PET SUVR image, and FreeSurfer brain mask files were identified for older participants and YCs. MR images underwent automated brain extraction with the Robust Learning-Based Brain Extraction System (ROBEX; https://www.nitrc.org/projects/robes)67 for linear and nonlinear transformation into common space (MN1152) with 2 mm voxel resampling. MR and SUVR images were registered to MN1152 space with the resulting nonlinear transformation.

Aligned SUVR images for YCs were merged into a four-dimensional matrix and the mean (μvux) and
standard deviation ($\sigma_{\text{vox}}$) were calculated for each voxel. Participant aligned SUVR images were then z-scored for each voxel (white square), the SUVR value is identified for the older participant. The SUVR values for the corresponding voxel are identified in all YC scans, which are then used to calculate mean and standard deviation. The older participant’s voxel z-score is next calculated from participant SUVR and YC mean and standard deviation. Voxels are then classified as normal or abnormal with the threshold $z > 1.96$. Once all voxels have been classified, a region of interest (ROI) mask is applied including the cortex, hippocampus, and amygdala. TSS is calculated as the proportion of abnormal voxels within the ROI mask. All abnormal voxels can be displayed overlaying the participant’s MRI.

Next, individualized brain masks from FreeSurfer were applied to the native z-score images selecting for cortical regions, the hippocampus, and the amygdala. Remaining voxels within this mask were then thresholded for statistical significance ($p \geq 1.96$) relative to YCs and binarized to evaluate whether the voxel has abnormal tau pathology.

TSS was finally calculated as the proportion of abnormal voxels in the brain relative to the total number of candidate voxels (TSS = $\frac{\# \text{Voxels Abnormal}}{\# \text{Voxels Total}}$).

Cortical surface projections

Vertex-wise maps were created to visualize tau intensity (Fig. 2a) and tau spread (Fig. 2b) across disease-stage groups using tau PET SUVR images and abnormality masks respectively. Using FSL, participant SUVR image and previously-calculated abnormality mask were co-registered to MN152 space in the same process as described for TSS and then voxel-wise averages were taken for each disease stage. Tau intensity is depicted as the average tau PET SUVR for each voxel. Tau spread is depicted as the proportion of participants with abnormal tau pathology in each voxel. The data was then projected onto the cortical surface with an fsaverage template and functions from the nilearn package (v0.10.1) in Python (v3.9.6).

Statistics

All analyses were conducted with R v4.1.0 with a p-value threshold of <0.05. Semi-nested linear models referred to as the Four Model Comparison Framework (FMCF) were implemented in these analyses to evaluate the predictive power of TI and TSS separately and conjointly for a variable (X), accounting for covariates age and sex, with the following format: Model 1 ($X \sim TI + \text{Covariates}$), Model 2 ($X \sim TSS + \text{Covariates}$), Model 3 ($X \sim TI + TSS + \text{Covariates}$), and Model 4 ($X \sim TI + TSS + TI \times TSS + \text{Covariates}$). Models 1 and 2 are therefore nested in Model 3 which is in turn nested in Model 4, however Models 1 and 2 are not nested within one another. The independent TI (Model 1) and independent TSS (Model 2) models assess the separate predictive ability of TI and TSS. The additive model (Model 3) assesses whether TI and TSS have unique predictive power or whether they provide redundant information with one metric as the better predictor. The interactive model (Model 4) assesses whether TI and TSS interact with one another in addition to independent effects, indicating a more complex and reliant relationship between the biological processes for tau intensity and tau spatial spread. Akaike Information Criterion (AIC) was used for model evaluation.
Disease-stage groups

Both tau metrics, TI and TSS, were first evaluated in their ability to discriminate between disease-stage groups. The Kruskal–Wallis test was conducted to determine significant difference in tau metric between disease-stage groups. Post-hoc comparisons were conducted between each group using Dunn’s test with Bonferroni p-value adjustment.

The FMCF was implemented with multinomial logistic regressions predicting disease-stage group to evaluate whether the tau metrics provide unique information for progression through the disease stages. Main effects and interaction were assessed with likelihood-ratio chi-square tests.

If an interaction was found, the relationship between TI and TSS was further dissected by direct comparison. We assessed the correlation between TI and TSS using Spearman correlation generally across all participants and additionally separated by disease-stage group. Changes in correlation across disease stages were assessed using Fisher’s Z Test. Relative amounts of TI and TSS were additionally compared for disease stages by calculating slopes from ranged major axis (RMA) regression to account for noise in the measurement of both TI and TSS then compared with 95% confidence interval.

Centiloid

TI and TSS were then evaluated relative to amyloid Centiloid as a proxy for time within AD in order to dissect the sensitivity of both tau metrics to within-group changes. We first validated the relationship between Centiloid and both tau metrics across all participants using Pearson correlation. The FMCF was additionally implemented with linear regression predicting Centiloid. Main effects and interaction were assessed with t-tests.

Within-group analyses were then conducted by calculating Pearson correlations between Centiloid and both tau metrics for each disease-stage group to determine which stages show elevated TI and TSS. We then evaluate whether later stages have elevated slope within-stage for TI and TSS, indicating acceleration of tau pathology later in AD, by evaluating the interaction between disease-stage group and Centiloid using F-tests with post-hoc analyses of the interaction using Tukey HSD.

TI and TSS were finally compared for their sensitivity to detecting change in tau pathology within AD after amyloid positivity. TI and TSS were log-transformed for normality then z-scored against the OC group in order to directly compare the tau metrics on the same scale. Local regression via LOESS was used to determine whether z-scored TI (zTI) and TSS (zTSS) visually diverge and at what point along the Centiloid scale. This divergence was then quantified by calculating Z-Score Difference (zTSS − zTI) within individuals and evaluating it against Centiloids with Pearson correlation for all participants and separated by disease-stage group. Sensitivity differences at later time points in AD were

Fig. 2: Surface projections for tau PET SUVRs and frequency of abnormality across disease stages. (Left) Voxel-wise tau PET SUVRs averaged across group. Color coded for SUVR (range 1–2). (Right) Voxel-wise group frequency of abnormality, determined by z-scoring against younger controls and thresholding for significance at +1.96. Color coded for the proportion of participants (range 0–1) in which the voxel is “abnormal.”
then evaluated by comparing the slopes of the disease-stage groups via interaction between Centiloid and disease-stage group using an F-test and post-hoc analyses of the interaction using Tukey HSD. It is important to note that sensitivity, as evaluated in these analyses, is relative to older controls who are amyloid negative, not based on a histopathology reference standard of tau pathology.

Role of funders
The study sponsors had no role in the study design, data collection, data analysis, data interpretation, writing of the report, or the decision to submit the manuscript for publication. All authors had full access to the data in the study and the corresponding author had final responsibility for the decision to submit for publication.

Results
Participants from Knight ADRC (n = 445) were included in the study and split into early Alzheimer disease stages. Of these individuals, 255 were classified as OC, 131 as preclinical AD, and 59 as symptomatic AD. 21 YCs were identified between Knight ADRC and DIAN. Participant disease-stage group and YC descriptive statistics are shown in Table 1.

Disease-stage groups
TI and TSS were first assessed as separate biomarkers for AD stage (Fig. 3a and b). Analyses of TI showed a significant main effect for disease stage (H(2) = 112.92, p < 0.0001). Post-hoc comparisons indicated that mean TI (Supplementary Table S1) was significantly different between all disease-stage groups. Analyses of TSS likewise showed a significant main effect for disease stage (H(2) = 91.96, p < 0.0001). Post-hoc comparisons indicated that mean TSS (Supplementary Table S2) was significantly different between all disease-stage groups. Both TI and TSS therefore demonstrate significantly elevated levels between all disease stages.

The FMCF was used to compare TI and TSS between disease stages (Table 2). ROC curves for each model are provided in Supplementary Figure S1. The independent models demonstrated that disease stage was predicted significantly by TI alone and by TSS alone. The additive model demonstrated that TI retained significance but TSS was not a significant predictor of diseases stage when modeled with TI. The interactive model showed no significant interaction between TI and TSS. The FMCF indicates higher predictive power of TI than TSS.

TI and TSS were moderately correlated overall (R = 0.79, p < 0.0001). Correlations were further assessed after splitting participants by disease-stage group (Fig. 3c). TI and TSS were moderately correlated for OC and preclinical AD groups and highly correlated for the symptomatic AD group. The symptomatic AD group (z = −2.75, p = 0.0060) and preclinical AD group (z = −2.25, p = 0.0244) showed significantly higher correlations between TI and TSS compared to the OC group, indicating that TI and TSS become better correlated after amyloid positivity. The relative levels of TI to TSS were further assessed for each disease-stage group (Supplementary Table S3). The symptomatic AD

<table>
<thead>
<tr>
<th>Younger control</th>
<th>Older control</th>
<th>Preclinical AD</th>
<th>Symptomatic AD</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>21</td>
<td>255</td>
<td>131</td>
</tr>
<tr>
<td>Age (years)</td>
<td>37.54 (10.06)</td>
<td>69.24 (7.98)</td>
<td>71.10 (7.24)</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>7 (33%)</td>
<td>128 (50%)</td>
<td>41 (31%)</td>
</tr>
<tr>
<td>Female</td>
<td>14 (67%)</td>
<td>127 (50%)</td>
<td>90 (69%)</td>
</tr>
<tr>
<td>MMSE</td>
<td>29.19 (1.03)</td>
<td>29.29 (1.07)</td>
<td>29.21 (1.20)</td>
</tr>
<tr>
<td>CDR</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>21 (100%)</td>
<td>255 (100%)</td>
<td>131 (100%)</td>
</tr>
<tr>
<td>0.5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Education (years)</td>
<td>16.28 (2.02)</td>
<td>16.50 (2.33)</td>
<td>16.36 (2.14)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>14 (67%)</td>
<td>223 (87%)</td>
<td>117 (89%)</td>
</tr>
<tr>
<td>Black</td>
<td>2 (10%)</td>
<td>30 (12%)</td>
<td>12 (9%)</td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>5 (24%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Participant sample size and characteristics for healthy controls and participants, split by disease-stage groups. Data are formatted as n (%) or mean (SD). AD, Alzheimer disease. MMSE, Mini-mental state examination. CDR, Clinical dementia rating.

Table 1: Participant demographics.
group was significantly different from the other two disease-stage groups, demonstrating a higher ratio of TI relative to TSS and indicating a change in relationship between TI and TSS after cognitive impairment is observed.

### Centiloid

Across all participants, TI ($R = 0.57$, $p < 0.0001$) and TSS ($R = 0.51$, $p < 0.0001$) were significantly larger at higher Centiloid values. The FMCF was once again used to compare TI and TSS relative to Centiloid (Table 3, Supplementary Tables S4 and S5). The independent models demonstrated significant main effects for both TI and TSS in predicting Centiloid when modeled separately. The additive model demonstrated that when TI and TSS are modeled together, TI retained significance but TSS was not significant. However, the interactive model showed a significant interaction between TI and TSS. TI and TSS are therefore strong predictors of Centiloid independently, but a more complex and interactive relationship is identified between TI and TSS across Centiloids.

Within-group analyses were conducted for TI and TSS across Centiloid. Fig. 4a shows a positive relationship between TI and Centiloid for the preclinical AD and symptomatic AD groups, but not for the OC group. The interaction between Centiloid and disease-stage group was highly significant (Supplementary Table S6), indicating further elevation in TI across Centiloids for later disease stages. Post-hoc pairwise analyses (Supplementary Tables S7 and S8) confirmed significant distinction in slope between OC and preclinical AD, preclinical AD and symptomatic AD, and OC and symptomatic AD groups. Fig. 4b shows a positive relationship between TSS and Centiloid for the preclinical AD and symptomatic AD groups, but not for the OC group. The interaction between Centiloid and disease-stage group was highly significant (Supplementary Table S9). Pairwise analyses (Supplementary Tables S10 and S11) confirmed significant distinction in slope between OC and symptomatic AD as well as between preclinical AD and symptomatic AD groups. However, unlike with TI, significance was not found between OC and preclinical AD groups.

### Table 2: Disease stage model comparison

<table>
<thead>
<tr>
<th>AIC</th>
<th>Tested variable</th>
<th>df</th>
<th>N</th>
<th>$\chi^2$</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Groups $\sim$ TI</td>
<td>644.70</td>
<td>2</td>
<td>445</td>
<td>172.78</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Groups $\sim$ TSS</td>
<td>649.90</td>
<td>2</td>
<td>445</td>
<td>122.58</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Groups $\sim$ TI + TSS</td>
<td>647.43</td>
<td>2</td>
<td>445</td>
<td>51.47</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Groups $\sim$ TI + TSS + TI $\times$ TSS</td>
<td>649.02</td>
<td>3</td>
<td>445</td>
<td>14.90</td>
<td>0.00026</td>
</tr>
<tr>
<td>Groups $\sim$ TI + TSS + TSS $\times$ TSS</td>
<td>644.14</td>
<td>2</td>
<td>445</td>
<td>1.27</td>
<td>0.39</td>
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<tr>
<td>Groups $\sim$ TI + TSS + TI $\times$ TSS</td>
<td>645.21</td>
<td>2</td>
<td>445</td>
<td>2.41</td>
<td>0.30</td>
</tr>
<tr>
<td>Groups $\sim$ TI + TSS + TSS $\times$ TSS</td>
<td>644.02</td>
<td>3</td>
<td>445</td>
<td>14.90</td>
<td>0.00026</td>
</tr>
<tr>
<td>Groups $\sim$ TI + TSS + TI $\times$ TSS</td>
<td>644.14</td>
<td>2</td>
<td>445</td>
<td>1.27</td>
<td>0.39</td>
</tr>
<tr>
<td>Groups $\sim$ TI + TSS + TSS $\times$ TSS</td>
<td>645.21</td>
<td>2</td>
<td>445</td>
<td>2.41</td>
<td>0.30</td>
</tr>
</tbody>
</table>

Comparison of individual TI, individual TSS, additive, and interactive models predicting disease-stage group. Multinomial logistic regressions with covariates age and sex evaluated with likelihood ratio chi-square tests. Test statistics and significance reported for tested variables as well as evaluation of the model via AIC. TI, Tau Index. TSS, Tau Spatial Spread. AIC, Akaike Information Criterion. df, degrees of freedom.

**Fig. 3:** Tau Index and Tau Spatial Spread are elevated across disease stages. (a-b) Comparison of Tau Index (a) and Tau Spatial Spread (b) across disease-stage groups. Pairwise comparisons conducted using Dunn’s test with Bonferroni p-value adjustment (c) Tau Index and Tau Spatial Spread plotted for each disease-stage group with ranged major axis regression and Spearman correlation reported with 95% confidence interval and p-value. Shaded area refers to 95% confidence interval of the regression slope.
indicating TI is more sensitive than TSS for differences observed between OC and preclinical AD groups.

A deviance was observed between \( z_{TI} \) and \( z_{TSS} \) relative to Centiloid at the point of amyloid-positivity,24–26 around 20–25 Centiloids (Fig. 4c). The Z-Score Difference was negatively correlated to Centiloids \( (R = -0.52, p < 0.0001) \) overall (Fig. 4d). However, this correlation was only maintained within preclinical AD \( (R = -0.33, p = 0.00011) \) and symptomatic AD \( (R = -0.32, p = 0.014) \) disease stages. The interaction between Centiloid and Group was significant (Supplementary Tables S12–S14). Z-score Difference was therefore elevated within and between disease stages, indicating TI is increasingly more sensitive than TSS throughout AD progression after amyloid positivity.

### Discussion

The progression of tau pathology in AD can be deconstructed into two key components: the spread of tau into new regions (tau spatial spread) and the simultaneous accumulation of NFTs in regions already impacted by tau (tau intensity). The characterization and quantification of these biological processes in tandem using neuroimaging is important for evaluating patient outcomes and predicting subsequent clinical and cognitive decline. The primary goal of these analyses was to propose a method for quantifying tau spatial spread to evaluate and compare tau spatial spread and tau intensity during the early stages of AD.

We found that both tau intensity and tau spread are elevated across all disease stages, beginning as early as the preclinical AD stage. Both tau metrics (TI and TSS) increased relative to amyloid indicating that they are sensitive measures across a continuous scale of disease progression. More tau spread is observed at earlier stages of AD, however tau intensity increases once individuals become symptomatic, at which point TI and TSS are highly correlated. TSS however demonstrates greater variability between subjects, resulting in greater predictive power of TI.

Previous neuroimaging research supports a temporal order of AD biomarkers in which tau pathology begins developing after amyloid positivity but prior to symptom onset,24,25 thereafter positively correlated with cognitive decline.23 Our results are consistent with this literature, demonstrating increasingly elevated levels of TI between disease stages, defined by amyloid-beta and cognitive status. We also found positive correlations to amyloid-beta within disease stages after a participant is amyloid-positive indicating that once tau pathology begins, there is an association with amyloid-beta. However, tau progression is not limited to intensity alone. A spatiotemporal order of brain regions impacted by tau pathology has been described following Braak staging simultaneous to the increasing accumulation of NFTs in early-impacted regions.19,21,75,76 Our results similarly demonstrate elevated TSS in addition to TI across disease stages and amyloid-beta burden. This suggests the general accelerated rate of tau pathology progression after cognitive impairment can be attributed to both increased spread and intensity.

Despite the observation of simultaneous tau spread and tau intensity, the biological mechanisms for these processes suggests a temporal order. Tau is believed to spread primarily via activity-dependent neuronal propagation,19 however increasing tau intensity with the development of NFTs disrupts neuronal transport and ultimately precipitates cell death.77 Tau spread therefore necessitates functional active neurons and must precede tau intensity. Consistent with this order, we found relatively higher levels of TSS than TI in the earlier disease stages representative of initial tau pathology development. The finding of early tau spread prior to substantial tau accumulation is supported by previous work in which widespread tau aggregates are found at early Braak stages.19 Once individuals become symptomatic, however, TI is elevated relative to TSS and the two metrics become strongly correlated. This could be explained by early tau spread impacting substantial regions by this point. The dynamic accumulation of NFTs within such regions ultimately results in an acceleration of tau intensity while tau spread is sustained.

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**Table 3: Centiloid model comparison.**

<table>
<thead>
<tr>
<th>AIC</th>
<th>Tested variable</th>
<th>( \beta )</th>
<th>Lower CL</th>
<th>Upper CL</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Centiloid ( \sim ) TI</td>
<td>3437.92</td>
<td>TI</td>
<td>20.50</td>
<td>17.53</td>
<td>23.48</td>
</tr>
<tr>
<td>Centiloid ( \sim ) TSS</td>
<td>3459.51</td>
<td>TSS</td>
<td>50.93</td>
<td>42.50</td>
<td>59.36</td>
</tr>
<tr>
<td>Centiloid ( \sim ) TI + TSS</td>
<td>3427.24</td>
<td>TI</td>
<td>16.64</td>
<td>11.45</td>
<td>21.74</td>
</tr>
<tr>
<td></td>
<td>─</td>
<td>TSS</td>
<td>12.53</td>
<td>-7.59</td>
<td>27.59</td>
</tr>
<tr>
<td>Centiloid ( \sim ) TSS + TI × TSS</td>
<td>3424.32</td>
<td>TI</td>
<td>23.65</td>
<td>17.18</td>
<td>30.12</td>
</tr>
<tr>
<td></td>
<td>─</td>
<td>TSS</td>
<td>55.65</td>
<td>29.25</td>
<td>82.05</td>
</tr>
<tr>
<td></td>
<td>─</td>
<td>TI × TSS</td>
<td>-24.76</td>
<td>-37.31</td>
<td>-12.22</td>
</tr>
</tbody>
</table>

Comparison of individual TI, individual TSS, additive, and interactive models predicting Centiloid. Linear regressions evaluated with covariates age and sex with t-tests. Beta coefficient estimates with 95% confidence level and significance reported for tested variables as well as evaluation of the model via AIC. TI, Tau Index. TSS, Tau Spatial Spread. AIC, Akaike Information Criterion. CL, Confidence Limit.
Fig. 4: Relationship of Tau Index and Tau Spatial Spread relative to amyloid burden. Tau metrics assessed against Centiloid. Shaded area refers to 95% confidence band. (a–b) TI and TSS for participants relative to Centiloid split by disease-stage group. Pearson correlation reported with 95% confidence interval and p-value. (c) Z-Scored TI and TSS relative to Centiloid with local regression. (d) Divergence of TI and TSS calculated by difference between z-scored TSS and z-scored TI and split by disease-stage group with Pearson correlation.
Evaluating TSS may therefore be important in some cases for identifying early changes in AD prior to cognitive decline. TSS could also be robust to atypical presentations of AD in which tau spread occurs along a different regional pattern than typical amnestic AD. While TI is constrained to key regions in typical AD, TSS can capture tau throughout the cortex. This difference between tau metrics is a strength of TSS. However, this lack of constraint also means TSS may capture erroneous tau PET signal or low levels of tau pathology that is not attributed to AD as demonstrated in the variability observed in the YC and OC groups (Supplementary Figures S2 and S3). It is important to note however that this variability is likewise observed in the TI metric, with a positive correlation to age potentially indicating Primary Age-Related Tauopathy (PART).

The calculation of TSS uses binary classification of whether a voxel is abnormal relative to younger controls. The threshold for each voxel is therefore relatively low and non-AD tau signal may surpass the threshold and classify voxels as abnormal. These false-positive voxels hold equal weight to voxels with much higher AD-related tau signal in the quantification of TSS, introducing potential noise and variability. TI, however, is an intensity-based approach that inherently accounts for the difference in strength of tau signal, minimizing such noise and variability, while additionally selectively including regions characteristic of AD-related tauopathy. TI may therefore be more specific to AD-related tauopathy, which corresponds to our observation of greater variability in TSS than TI within disease stages. This also explains why we find TI is more sensitive than TSS to differences between disease stages.

TI and TSS therefore have complementary strengths and weaknesses and should be used in conjunction for stronger analyses. Individuals with widespread false positives such as those exhibiting diffuse cortical uptake have additionally been identified by comparing relative TI and TSS values (Supplementary Figures S4 and S5).

Due to largely unsuccessful results with amyloid-targeting drugs, many clinical trials have changed focus to participants with preclinical AD to prevent or delay symptom onset. New tau-targeting drugs have additionally shown promise in animal models and are currently in early clinical trial phases. With the surge of these trials, evaluating both TI and TSS may provide additional insight into the success of tau-targeting drugs in reducing or slowing tau progression, particularly in the preclinical stage at which the metrics show the largest divergence.

In interpreting the results of our study, limitations should be considered. The Knight ADRC dataset is skewed towards cognitively normal participants in order to capture early AD-related changes. A longitudinal study with multiple time points of tau PET with participants who progress from preclinical to symptomatic AD will be an important follow up study. There is also a strong ascertainment bias in which any group of individuals who participate in longitudinal imaging and biomarker studies are not representative of the general population, so these results may not generalize. The limited sample size does not provide the power for analyses evaluating potential confounding factors outside of those included in this paper, which may influence the results and interpretation of this study. Some statistical models are additionally simplified for interpretability, such as the usage of linear regression within the FMCF predicting Centiloid despite violations of the underlying assumptions. Subsequent analyses for the tested relationships support the reported findings despite such violations.

It is important to note that TI and TSS differ in both the method of quantification and the regions assessed. TI is restricted to early regions of interest while TSS accounts for many regions throughout the brain in order to determine whether there is added benefit in evaluating tau spread into additional brain regions. Restricting the regions included in TSS or expanding upon the regions included in TI would allow for more specific analyses between the two metrics, but would likewise reduce the interpretability of the tau metrics regarding their corresponding components of tau pathological progression.

TSS could additionally implement alternative approaches to classify tau positivity at the voxel level such as Gaussian mixture modeling. However, we expect to see a continuum of tau PET SUVRs at the voxel level rather than a clear separation between individuals who are tau-negative and tau-positive since the included cohort encompasses early stages of AD and therefore relatively low levels of tau pathology. Gaussian mixture modeling and similar methods would therefore choose an arbitrary cutoff for tau positivity and be extremely computationally expensive at a voxel-level. We instead chose to classify tau positivity based on a z-score threshold of 1.96 relative to young controls, who are not expected to have tau pathology, because this method identifies when a voxel shows statistically significant SUVRs and therefore abnormally high levels of tau.

Several study strengths result from the chosen method of calculation for TSS. In this study, “abnormal” voxels are identified based on relative SUVR values compared to a younger control group. This method accounts for regional variability in SUVRs not attributed to tau pathology and therefore reduces voxel-wise false positives. When calculating the final value for TSS, an ROI mask is additionally applied to the tau PET scan in order to select for biologically-relevant regions of interest including cortical ROIs, the hippocampus, and the amygdala. The implementation of the ROI mask filters out false positives from off-target PET tracer binding. With the reduction of false positives, TSS better represents AD tau pathology spread extent.
Our results reveal a complex pattern of tau progression in early AD in which tau spatial spread and tau intensity demonstrate different temporal patterns. Tau spatial spread may be captured earlier than tau intensity but shows subject-specific variability that tau intensity is less vulnerable to. Tau intensity increases more rapidly relative to tau spatial spread upon symptom onset and at this point spread and intensity are highly correlated. These results suggest a critical preclinical period in which tau spread and tau intensity behave differently which could have broad applications in preventative tau-targeting clinical trials.

Contributors
SD developed the programming scripts, analyzed the data, generated the figures, conducted the literature search, and wrote the manuscript. SD, AM, BAG, and TLSB contributed to method conceptualization. SD, AM, BAG, CDC, NM, DH, and CX developed the statistical approach. SD, AM, BAG, CDC, NM, DH, SK, SF, JS, HS, SJ, KJ, RCH, BMA, CX, AJA, JH, CC, AD, RJ, JCM, and TLSB contributed to data interpretation. TLSB, SK, SF, JS, HS, AD, CC, and RCH oversaw data quality control and processing. TLSB, RJB, AD, CC, and JCM oversaw overall study design and general implementation. AJA, JH, TLSB, RJB, JCM, SJ, AD, CC, and KJ oversaw study implementation and data collection. SD, AM, BAG, CDC, NM, DH, SK, SF, JS, HS, SJ, KJ, RCH, BMA, CX, AJA, JH, CC, AD, RJ, JCM, and TLSB revised the manuscript. This manuscript has been reviewed by DIAN study investigators for scientific content and for consistency of data interpretation with previous DIAN study publications. DIAN investigators oversaw the collection of all demographic, clinical, neuroimaging, and genetic underlying data for participants chosen from the DIAN-Observational study. SD, AM, BAG, and TLSB accessed and verified all included DIAN data. Underlying data from the Knight ADRC was accessed and verified by several authors for included demographic (SD, AM, BAG, TLSB), clinical (SD, AM, BAG, TLSB), and neuroimaging (SD, AM, BAG, CDC, SK, SF, JS, HS, RCH) data. Avid Radiopharmaceuticals, Inc., a wholly owned subsidiary of Eli Lilly and Company, enabled use of the 18F-flortaucipir tracer by providing precursor, but did not provide direct funding and was not involved in data analysis or interpretation. All authors contributed substantially to the conception or design of the work or the acquisition of data for the work. All authors reviewed and approved the final manuscript and agree to be accountable for all aspects of the work.

Data sharing statement
The data used in these analyses is available upon request for Knight ADRC at https://knightheadrc.wustl.edu/data-request-form/ and for DIAN at https://dian.wustl.edu/our-research/for-investigators/dian-observational-study-investigator-resources/data-request-form/.

Declarations of interests
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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.ehram.2024.105080.

References
