Plasma p tau 181 are associated with white matter microstructural changes in Alzheimer’s disease

Fardin Nabizadeh (fardinnabizade1378@gmail.com)  
Neuroscience Research Group (NRG), Universal Scientific Education and Research Network (USERN), Tehran, Iran

Seyed Behnamedin Jameie  
Neuroscience Research Center (NRC), Iran University of Medical Science, Tehran, Iran

Saghar Khani  
Iran university of Medical science

Aida Rezaei  
School of Medicine, Tehran University of Medical Science, Tehran, Iran

Niloofer Deravi  
Student Research committee, School of medicine, Shahid Beheshti University of Medical Science, Tehran, Iran

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Abstract

Alzheimer's Disease (AD) is characterized by cognitive impairments and memory difficulties, which cause daily activities, personal and behavioural problems. In recent years blood-based biomarkers like plasma phosphorylated tau protein at threonine 181 (p tau 181) emerged as new tools and showed a sufficient power in detecting AD patients from healthy people. Here we investigate the correlation between p tau 181 and white matter microstructural changes in AD. We add 41 Alzheimer diagnosed patients, 155 participants with mild cognitive impairments and 43 healthy controls with baseline plasma p tau 181 level and DTI values for each region of interest from the ADNI database. The analysis revealed that the plasma level of p tau 181 could predict changes of MD (Mean Diffusivity), RD (Radial Diffusivity), DA (Axial Diffusivity) and FA (Fractional Anisotropy) parameters in widespread regions and there is a significant association between white matter pathways alteration in different regions and p tau 181 plasma measurement within each group. In conclusion, our findings showed that plasma p tau 181 levels are associated with cellular and molecular changes in AD, which enhance this biomarker's for diagnostic procedures and support the application of plasma p tau 181 as a biomarker for white matter changes and neurodegeneration. Longitudinal studies are also necessary to prove the efficacy of these biomarkers and predicting role in structural changes.

Introduction

Alzheimer's disease (AD) is the course of dementia and memory deficits that affect millions of people and responsible for cognitive and functional decline, mostly in aged people (1, 2). AD is characterized by cognitive impairments and memory difficulties, which cause daily activities, personal and behavioural problems (1, 3). AD is associated with the formation of neurofibrillary tangles (NFTs), including hyperphosphorylated tau protein (p tau) and extracellular Amyloid β (Aβ) plaques in regions responsible for memory and other cognitive functions such as hippocampus (4, 5). NFTs are typical brain lesions consist of aggregated and hyperphosphorylated forms of tau protein, which leads to loss of its ability to binding microtubules and assembled into paired helical filaments. Intracellular NFTs in regions involved in cognitive functions are associated with cognitive decline by the disruption in axonal transport and neural loss (6, 7).

Many researchers reported Aβ, total Tau and Tau phosphorylated at threonine 181 (p tau 181) in cerebrospinal fluid (CSF), and Positron emission tomography (PET) as biomarkers for Alzheimer diagnoses. However, in recent years plasma biomarkers emerged as new diagnostic tools and showed sufficient efficacy in detecting AD patients from healthy people (8). Blood-based measures showed that p tau 181 might be a reliable biomarker for Alzheimer's and disease progression (9-11). However, there is no evidence that shows the effect of plasma p tau 181 on white matter connections and neurodegeneration in the brain of AD patients. We hypothesized plasma p tau 181 level could predict structural brain connections changes in regions play a significant role in cognitive, learning and memory function. In the
present research, we investigated the correlation between p tau 181 in serum and diffusion tensor imaging (DTI) values in an observational cross-sectional study to address this question.

**Materials And Methods**

**Data Acquisition**

Participants information acquired from the Alzheimer's Disease Neuroimaging Initiative (ADNI) database (adni.loni.usc.edu). The ADNI leading by Principal Investigator Michael W. Weiner, MD was launched in 2003 as a public-private partnership. The primary goal of ADNI has been to test whether serial magnetic resonance imaging (MRI), positron emission tomography (PET), other biological markers, and clinical and neuropsychological assessment can be combined to measure the progression of mild cognitive impairment (MCI) and early Alzheimer's disease (AD). We extracted baseline visit data for whom demographic data, post-processed DTI, CSF Amyloid β, CSF p tau, CSF total, and plasma p tau 181 tau levels from baseline visit were available. We also downloaded results of white matter hyperintensity analysis and brief volumetric data for better understanding. Our cross-sectional study consisted of 21 patients with AD, with their baseline plasma p tau 181, CSF Amyloid β, CSF total tau, and CSF p tau levels and their post-processed Diffusion tensor imaging (DTI).

Participants classified as AD patients if their mini-mental state examination (MMSE) was between 20 and 24, clinical dementia rating (CDR) score between 0.5 and 1, and met the National Institute of Neurologic and Communicative Disorders and Stroke and the Alzheimer's Disease and Related Disorders Association criteria(12).

**Plasma p tau 181 Measurements**

Plasma samples analyzed at University of Gothenburg, Sweden by using Single-Molecule array (Simoa) technique. The detailed procedure described in (adni.loni.usc.edu). (11).

**DTI Imaging Processing**

We extracted the results of DTI ROI analysis from ADNI. For each subject, All images were corrected, extracerebral tissue removed, and normalized.

Extraction Tool (BET) from FSL (13). To align data from different subjects into the same 3D coordinate space, each T1-weighted anatomical image was linearly aligned to a version of the Colins27 brain template (14) using FSL's flirt (15) with 6 degrees of freedom to allow translations and rotations in 3D. The Colin27 brain was zero-padded to have a cubic isotropic image size 220x220x220 1mm^3) and then downsampled (110x110x110 2mm^3) to be more similar to the DWI resolution.

To correct for echo-planar imaging (EPI) induced susceptibility artifacts, which can cause distortions at tissue-fluid interfaces, skull-stripped b0 images were linearly aligned to their respective T1-weighted structural scans using FSL's flirt with 9 degrees of freedom and then elastically registered to their aligned
T1 scans using an inverse consistent registration algorithm with a mutual information cost function (16) as described in (17). The resulting 3D deformation fields were then applied to the remaining 41 DWI volumes before mapping diffusion parameters. To account for linearly registering the average b0 from the DWI images to the structural T1-weighted scan, a corrected gradient table was calculated.

A single diffusion tensor was modeled at each voxel in the brain from the eddy- and EPI-corrected DWI scans using FSL's \textit{dtifit} command, and scalar anisotropy and diffusivity maps were obtained from the resulting diffusion tensor eigenvalues ($\lambda_1, \lambda_2, \lambda_3$). Fractional anisotropy (FA) was calculated from the standard formula.

We registered the FA image from the JHU DTI atlas (18) to each subject using a previously described mutual information-based elastic registration algorithm(16). We then applied the deformation to the stereotaxic JHU “Eve” WM atlas labels (http://cmrm.med.jhmi.edu/cmrm/atlas/human_data/file/AtlasExplanation2.htm) using nearest neighbour interpolation to avoid intermixing of labels.

This placed the atlas ROIs in the same coordinate space as our DTI maps. We were then able to calculate the average FA and MD within the boundaries of each of the ROI masks for each subject. Of the 56 WM ROIs, we excluded 4 ROIs, the left and right middle cerebellar peduncle, and the pontine crossing tract, as they often fall wholly or partially out of the field of view (FOV). We note that this is also occasionally true of the left and right medial lemniscus, inferior and superior peduncles. We only included non-zero voxels within the FOV in our calculations of mean FA and MD. In addition to the 52 JHU labels, five more ROIs were evaluated: the bilateral fornix, bilateral genu, bilateral body, and bilateral splenium of the corpus callosum and the full corpus callosum, to get full summary measures of the regions.

Tensor based spatial statistics (19) was also performed, and the mean FA in regions of interest along the skeleton was extracted. TBSS was performed according to protocols outlined by the ENIGMA-DTI group: http://enigma.loni.ucla.edu/wpcontent/uploads/2012/06/ENIGMA_TBSS_protocol.pdf.

In short, all subjects were registered to the ENIGMA-DTI template in ICBM space, and standard \textit{tbss} steps were performed to project individual FA maps onto the skeletonized ENIGMA-DTI template. ROI extraction was also performed according to the following protocol to extract the mean FA in ROIs along with the skeleton: http://enigma.loni.ucla.edu/wpcontent/uploads/2012/06/ENIGMA_ROI_protocol.pdf.

**Cognitive assessments**

The patients' cognitive condition assessed by the Mini-Mental State Exam (MMSE) which is a common test of cognitive function among the elderly; it includes tests of orientation, attention, memory, language, and visual-spatial skills. MMSE scores were extracted for each patient from the ADNI Mini-Mental Examination.

**Statistical Analyses**
We used SPSS16 software for statistical analyses. First we tested normality of variables by Kolmogorov Smirnov and Shapiro Wilk tests. Then we log-transformed the nonnormal variables to normal distribution. We analyzed the difference between groups by one way ANOVA using Bonferroni correction for multiple comparisons. Then we used a Partial correlation adjusted for age, sex, and APOE for checking the relation between plasma p tau 181 and other demographic variables once among all participants and then within groups. We used the same model for investigating the association between plasma p tau 181 and DTI values in each ROI. The bootstrap method was used for addressing type I error due to multiple comparisons in correlation models.

Result

Patient characteristic

In this study, baseline cohort information of 203 participants was entered. The mean age was 73, including 117 men and 86 women. Participants were all educated, and the average length of the education was 15.9 years. Among all 102 participants had at least one APOE ε4. the mean scores of the MMSE test were 27.25. details of demographic information for each group described in Table1.

Between groups analysis

the level of plasma p tau 181 is significantly different between groups [F(2, 200)=9.05, p=0.000]. there is no meaningful different between group’s age and mean education time [F(2, 200)=0.425, p=0.655], [F(2, 200)=2.50, p=0.084]. the comparison of APOE genotype status between groups revealed a strong different [F(2, 200)=9.06, p=0.000], and the glucose uptake significantly lower in AD group [F(2, 200)=43.9, p=0.000].

plasma p tau 181 with characteristics

The Partial correlation model for p tau 181 level and each demographic variables, while adjusted for age, sex, and APOE genotype showed plasma p tau 181 has a strong correlation with glucose uptake in angular, temporal, and posterior cingulate [correlation coefficient:-0.231, p: 0.001], education time [correlation coefficient:-0.166, p: 0.019], MMSE score [correlation coefficient:-0.216, p: 0.002], and also there is a positive correlation between age and p tau while controlled for sex and APOE [correlation coefficient: 0.215, p: 0.002] among all participants. When we investigated the correlation between p tau 181 and demographic features within groups, the correlations started to disappearing, and there is only one correlation between p tau and glucose uptake in AD patients [correlation coefficient:-0.397, p: 0.017].

Plasma p tau 181 with microstructural changes

In the first, we assessed the relation between p tau level and DTI values in each ROI among all participants by implementing partial correlation model adjusted for effects of age, sex, and APOE genotype. There is a strong correlation between plasma measures of p tau 181 with a widespread pattern of changes in each FA, DA, RD, and MD in the brain (Tab2). We found a negative correlation between p
tau and FA in Left hippocampal cingulum, Splenium of right corpus callosum, Left and Right Tapatum. Moreover higher level of p tau 181 correlate with overall higher MD and DA in Splenium of right and left corpus callosum, Left and Right Tapatum, Right Posterior corona radiate, Right Sagittal stratum, Right Uncinate fasciculus, Retrolenticular part of right internal capsule, and Left Medial lemniscus (Tab2). Also, the correlation was same for RD in Left hippocampal cingulum, Splenium of right corpus callosum, Left and Right Tapatum, Right Posterior corona radiate, Right Sagittal stratum, Right Uncinate fasciculus, Retrolenticular part of right internal capsule, and Left Medial lemniscus (Tab2).

The investigating relation between p tau 181 and connectometry values within each group showed there is a significant correlation for MD, RD, and DA in Left Medial lemniscus, MD and DA in Left Inferior cerebellar peduncle, MD and RD in Right Superior cerebellar peduncle among AD patients (Tab3). The p tau correlates with MD and RD only in Left Medial lemniscus within MCI group (Tab3). Partial correlation revealed there is a correlation between p tau 181 and DA in Right Superior cerebellar peduncle and RD in Left Uncinate fasciculus among healthy controls (Tab3).

**Discussion**

In a cross-sectional study based on the ADNI cohort, we concluded that the plasma level of p tau 181 independently predicts microstructural changes in the brain of Alzheimer's patients. We used a step-by-step strategy; first, we investigate the difference between participants demographic, including age, glucose uptake, MMSE scores, education, APOE genotype, sex, and plasma p tau 181. Then we narrowed the study to examine the correlation between p tau 181 with demographic variables among all participants and within groups separately. Then we used a partial correlation model controlled for age, APOE and sex to investigate the relation between p tau 181 in plasma and changes in white matter. Baseline plasma p tau 181 levels are associated with widespread white matter changes in all participants in the disease's pathological signatures areas, including the hippocampal cingulum, Splenium of corpus callosum, Tapatum, Posterior corona radiate, Sagittal stratum, Uncinate fasciculus, Retrolenticular part of internal capsule, cerebellar peduncles, and Medial lemniscus. overall, according to observations, a decrease in FA and increase in MD reflect demyelination and axonal loss (20). this is the first study that investigated the relationship between plasma p tau 181 and white matter changes.

In the onset of dementia and cognitive decline, many areas seem to change. The internal capsule, corona radiate, Uncinate fasciculus, cerebellar peduncles, Medial lemniscus, and hippocampal cingulum in AD people change compared to healthy people without cognitive problems (21). In our study, plasma p tau 181 levels were significantly associated with these areas, indicating that our findings are in line with previous studies investigating microstructural changes related to Alzheimer's. As results of our analysis, plasma p tau 181 predicted structural changes in the left hippocampal cingulum, which indicate the importance of the cingulum as an important area in the pathological course of the disease, and there is also evidence of a link between CSF p tau and Aβ with a change of MD in the cingulum region (22). Moreover, the research results provide compelling evidence in support of our findings and state that brain connectivity in the posterior cingulum can be a good predictor for cognitive decline in Alzheimer's disease.
The cingulum bundle is an important white matter tract that connects the frontal, parietal, and medial temporal, linking the subcortical nucleus to the cingulate gyrus and extending into the hippocampal and parahippocampal regions, and for that damage in areas close to the hippocampus in the cingulum causes cognitive problems in many domains such as language, memory and executive control (24).

Previously CSF biomarkers revealed as a predictor for changes in the Uncinate fasciculus (25) and these regions involved in language processing and damage to uncinate fasciculus may cause language impairments (26) and as well as there is a significant correlation between plasma p tau 181 and changes in this region from our results. Similarly, X Li et al. Found that pathological levels of Aβ42 and CSF total tau in people with Alzheimer's-related cognitive impairments correlated with decreased FA and increased MD in the white matter pathway (27).

Biomarkers in CSF may be active several years before the onset of symptoms, and Aβ and Tau are most critical (28). Many studies have emphasized the diagnostic role of t tau and p tau in CSF and state that they can predict the progression to dementia (29-32). On the other hand, some studies present different findings (33, 34). Despite this, in predicting AD by each of these biomarkers alone, p-tau preferred in terms of specificity and sensitivity (28).

Studies have been conducted on blood-based biomarkers in recent years as a noninvasive and accessible marker for monitoring people with risk of developing Alzheimer. Evidence revealed that plasma p tau 181 and t tau levels have a high diagnostic value, and their levels are much higher in AD subjects than in MCI and healthy control (35). However, studies have described p tau 181 better than t tau (9, 10). The largest plasma p tau 181 study in the diagnosis of Alzheimer's, which included the results of four independent cohorts, states that plasma p tau 181 has a high performance in identifying the clinical diagnosis of Alzheimer's patients with an unknown amyloid status and was able to differentiate Alzheimer's disease from other neurodegenerative diseases and with Aβ PET can detect Alzheimer's in the early stages (11).

In conclusion, our study results show that plasma p tau 181 levels are associated with neurodegeneration in pathogenesis regions of Alzheimer's disease, which enhance this biomarker's diagnostic status and support the application of blood-based biomarkers as an early detector for white matter damages. Due to the increasing population with Alzheimer's and the resulting social costs and considering that AD's pathogenesis exists several years before its clinical signs, achieving a reliable biomarker with adequate sensitivity and specificity is necessary. Although plasma p tau 181 is superior to CSF biomarkers and imaging techniques in terms of availability, low cost, and non-invasiveness, More efforts should be made to standardize biomarkers' measurement and define their pathological threshold. Longitudinal studies are also necessary to prove the efficacy of these biomarkers predicting role in structural changes.

**Declarations**

*Data used in the preparation of this article were obtained from the Alzheimer's disease Neuroimaging Initiative (ADNI) database (adni.loni.usc.edu). As such, the investigators within the ADNI contributed to
the design and implementation of ADNI and/or provided data but did not participate in analysis or writing of this report. A complete listing of ADNI investigators can be found at:


**Corresponding Author:**

Nabizadeh F, Neuroscience Research Group (NRG), Universal Scientific Education and Research Network (USERN), Iran University of Medical sciences Tehran, Iran. E-mail: fardinnabizade1378@gmail.com

We have no conflicts of interest to disclose.

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**References**


Tables

**TABLE1.** Participants characteristic

<table>
<thead>
<tr>
<th>Demographic and health characteristic</th>
<th>CN(43)</th>
<th>MCI(119)</th>
<th>AD(41)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>72.9(±6.2)</td>
<td>72.8(±6.8)</td>
<td>74.0 (±8.6)</td>
<td>0.655</td>
</tr>
<tr>
<td>Sex(M/F)</td>
<td>22/21</td>
<td>71/48</td>
<td>24/17</td>
<td>0.625</td>
</tr>
<tr>
<td>Education, years</td>
<td>16.4(±2.6)</td>
<td>16.0(2.6)</td>
<td>15.2(±2.9)</td>
<td>0.084</td>
</tr>
<tr>
<td>MMSE</td>
<td>28.8(±1.4)</td>
<td>27.9(±1.9)</td>
<td>23.4(±1.8)</td>
<td>0.000</td>
</tr>
<tr>
<td>FDG-PET</td>
<td>1.30(±0.14)</td>
<td>1.29(±0.12)</td>
<td>1.06(±0.16)</td>
<td>0.000</td>
</tr>
<tr>
<td>APOE genotype</td>
<td>0.000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>With out ε4</td>
<td>31</td>
<td>58</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>One ε4</td>
<td>12</td>
<td>49</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>Two ε4</td>
<td>0</td>
<td>12</td>
<td>6</td>
<td></td>
</tr>
</tbody>
</table>

Values are showed as mean(±SD), Mini Mental State Examination(MMSE), fluorodeoxyglucose (FDG)-positron emission tomography (PET), results of ANOVA analysis between groups noted as p value

**Table2.** Significant Results of partial Correlation Analyses of DTI metrics and CSF P tau 181 Levels among all participants
<table>
<thead>
<tr>
<th>regions</th>
<th>Fractional anisotropy</th>
<th>Mean diffusivity</th>
<th>Radial diffusivity</th>
<th>Axial diffusivity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Correlation Coefficient</td>
<td>Correlation Coefficient</td>
<td>Correlation Coefficient</td>
<td>Correlation Coefficient</td>
</tr>
<tr>
<td>Left hippocampal cingulum</td>
<td>-0.218**</td>
<td>0.164</td>
<td>0.182*</td>
<td>0.124</td>
</tr>
<tr>
<td>Splenium of right corpus callosum</td>
<td>-0.142*</td>
<td>0.171*</td>
<td>0.168*</td>
<td>0.163*</td>
</tr>
<tr>
<td>Splenium of left corpus callosum</td>
<td>-0.101</td>
<td>0.141*</td>
<td>0.133</td>
<td>0.152*</td>
</tr>
<tr>
<td>Left Tapatum</td>
<td>-0.167*</td>
<td>0.166*</td>
<td>0.169*</td>
<td>0.156*</td>
</tr>
<tr>
<td>Right Tapatum</td>
<td>-0.175*</td>
<td>0.189**</td>
<td>0.194**</td>
<td>0.174*</td>
</tr>
<tr>
<td>Left Medial lemniscus</td>
<td>0.048</td>
<td>-0.178*</td>
<td>-0.163*</td>
<td>-0.174*</td>
</tr>
<tr>
<td>Left Posterior corona radiata</td>
<td>0.069</td>
<td>0.123</td>
<td>0.102</td>
<td>0.139</td>
</tr>
<tr>
<td>Right Posterior corona radiata</td>
<td>0.074</td>
<td>0.188**</td>
<td>0.147*</td>
<td>0.223**</td>
</tr>
<tr>
<td>Right Sagittal stratum</td>
<td>-0.002</td>
<td>0.181*</td>
<td>0.164*</td>
<td>0.199**</td>
</tr>
<tr>
<td>Right Uncinate fasciculus</td>
<td>-0.134</td>
<td>0.199*</td>
<td>0.202**</td>
<td>0.188**</td>
</tr>
<tr>
<td>Retrolenticular part of right internal capsule</td>
<td>-0.030</td>
<td>0.192**</td>
<td>0.178*</td>
<td>0.189**</td>
</tr>
</tbody>
</table>

*p < 0.05

**p<0.01

***p<0.001

Partial correlation coefficient of DTI metrics value of the brain regions and plasma p tau 181 levels controlled for age, APOE, and sex.