Longitudinal Cerebral Blood Flow and Amyloid Deposition: An Emerging Pattern?

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Although cerebral amyloid deposition may precede cognitive impairment by decades, the relationship between amyloid deposition and longitudinal change in neuronal function has not, to our knowledge, been studied. The aim of this article was to determine whether individuals without dementia with high and low amyloid burden show different patterns of longitudinal regional cerebral blood flow (rCBF) changes in the years preceding measurement of amyloid deposition. Methods: Twenty-eight participants without dementia (mean age ± SD, 82.5 ± 4.8 y; mildly impaired) from the Baltimore Longitudinal Study of Aging underwent yearly resting-state 15O-H2O PET scans for up to 8 y. 11C-PIB images of amyloid deposition were acquired on average 10.8 ± 0.8 y after the first CBF scan. 11C-PIB distribution volume ratios of regions of interest were estimated by fitting a reference-tissue model to the measured time–activity curves. On the basis of mean cortical distribution volume ratios, participants were divided into groups with high or low 11C-PIB retention. Differences in longitudinal rCBF changes between high- and low-11C-PIB groups were investigated by voxel-based analysis. Results: Longitudinal rCBF changes differed significantly between high- (n = 10) and low- (n = 18) 11C-PIB groups (P < 0.001). Greater longitudinal decreases in rCBF in the high-11C-PIB group than in the low-11C-PIB group were seen in right anterior to middle cingulate, right supramarginal gyrus, left thalamus, and midbrain bilaterally. Greater increases in rCBF over time in the high-11C-PIB group were found in left medial and inferior frontal gyri, right precuneus, left inferior parietal lobule, and left postcentral gyrus. Conclusion: In this group of older adults without dementia, those with high 11C-PIB show greater longitudinal decline in rCBF in certain areas, representing regions with greater decrements in neuronal function. Greater longitudinal increases in rCBF are also observed in those with higher amyloid load and may represent an attempt to preserve neuronal function in these regions.

Key Words: PIB; CBF; longitudinal; positron emission tomography; aging

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The development of in vivo amyloid imaging agents provides the opportunity to investigate prospective changes in amyloid deposition as a potential early marker of cognitive impairment and Alzheimer’s disease (AD) (1). Imaging studies have shown increased amyloid deposition in patients with AD and in some individuals with mild cognitive impairment (MCI), a condition thought to represent a transition between normal aging and AD (2). Studies of individuals without dementia focus on the ability of 11C-PIB and other amyloid imaging agents to differentiate between individuals who will and who will not progress to AD (1,3–6). One challenge in the use of amyloid imaging in early identification of individuals at risk for cognitive impairment and AD is that as many as 10%–30% of individuals who are clinically healthy on cognitive testing show amyloid deposition in imaging (7) and postmortem studies (7,8).

Evaluation of neuronal activity, using 15O-H2O, in conjunction with imaging of amyloid deposition, might help clarify the significance of amyloid deposition in older adults without dementia. In view of the suggestion by Buckner et al. (9) that default brain activity in young adulthood may affect amyloid deposition later in life, in this study we evaluated changes in resting-state regional cerebral blood flow (rCBF) that occur during the decade before measurement of amyloid deposition in older individuals without dementia. Such examination is important not only for improved understanding of the longitudinal pathophysiological relationship between rCBF and amyloid deposition but also because of potential clinical implications, as joint analysis of rCBF and 11C-PIB data may provide more diagnostic information than a single method alone.

As β-amyloid is associated with decreased resting rCBF in animal models (10), we hypothesized that individuals with higher amyloid deposition detected by 11C-PIB would exhibit different patterns of longitudinal rCBF change than those individuals with lower amyloid deposition. We predicted that the longitudinal changes in rCBF in subjects...
with high $^{11}$C-PIB retention would involve regions that show decreases in rCBF in individuals without dementia who eventually progress to AD. These regions would include the anterior and posterior cingulate, insula, middle and superior frontal, and superior temporal regions as well as the caudate nucleus and thalamus (11). In addition, we expected that at least some of the areas of change in longitudinal rCBF would spatially overlap regions of known high $^{11}$C-PIB retention, as vessel narrowing and subsequent hypoperfusion related to amyloid angiopathy may occur in some regions of $^{11}$C-PIB retention (12).

**MATERIALS AND METHODS**

**Study Participants**

Data from 28 participants without dementia (6 with a score of 0.5 on the Clinical Dementia Rating [CDR] (13) Scale) from the neuroimaging substudy (14) of the Baltimore Longitudinal Study of Aging who underwent both CBF and $^{11}$C-PIB scans were included in the study. On the basis of the degree of amyloid deposition as determined by the mean cortical distribution volume ratios (DVRs) described later, participants were divided into groups with high $^{11}$C-PIB retention and low $^{11}$C-PIB retention. Ten individuals (7 men and 3 women; mean age, 82.82 ± 5.11 y) had low $^{11}$C-PIB retention. High–and low–$^{11}$C-PIB-retention groups did not differ significantly with respect to age, sex, race, and education (Table 1). All participants underwent neuropsychological evaluation in conjuction with each neuroimaging visit. The CDR Scale, typically informant-based, was administered in conjuction with the $^{11}$C-PIB imaging study and was previously administered in conjuction with rCBF studies when participants scored 3 or greater on the Blessed-Information-Memory-Concentration (15) test. Cognitive status was determined by consensus diagnosis according to established procedures (8,16). A consensus diagnosis of MCI required progressive memory impairment without functional loss in activities of daily living (2). Informed consent was obtained from each participant before each study. Institutional Review Board approval was obtained for the study.

**$^{15}$O-H$_2$O Studies**

$^{15}$O-H$_2$O studies were performed annually for up to 9 assessments over an 8-y follow-up. During each imaging session, a resting PET scan was performed, during which the participants were instructed to keep their eyes open and focused on a computer screen covered by a black cloth. Scans were acquired on a scanner (4096+; GE Healthcare) with 15 slices (axial resolution, 6.5-mm full width at half maximum) and a 9-cm axial field of view. The superiormost portions of the parietal cortex were not evaluated because of the limited axial field of view. Images were obtained for 60 s once the total radioactivity counts in the brain reached threshold levels. A transmission scan in 2-dimensional mode using

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<tr>
<td>No. of participants</td>
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<td>CBF studies/participant</td>
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<td>Age at time of $^{11}$C-PIB study (y)</td>
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<td>Age range at time of $^{11}$C-PIB study (y)</td>
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<td>Sex (M/F)</td>
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<td>Education (y)</td>
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<td>White matter disease burden</td>
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<td>CDR status (no. with CDR = 0.5) at time of $^{11}$C-PIB study</td>
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<td>Diagnosis of MCI at time of $^{11}$C-PIB study</td>
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<td>Mean cortical DVR</td>
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*Fisher exact test for independence.
†MMSE results from 1 individual (CDR = 0) unavailable because of severe hearing impairment.
‡White matter ratings used 10-point scale, ranging from 0 (normal) to 9 (thick rind of periventricular white matter disease together with marked subcortical disease), from Cardiovascular Health Study (37,38).
§Exact Wilcoxon 2-sample test.
*Mean cortical DVR across orbital frontal, prefrontal superior frontal, lateral temporal, occipital, parietal, anterior cingulate, and posterior cingulate regions.

Results expressed as mean ± SD, with range in parentheses. Unless otherwise noted, results are based on t tests for independent samples. NS = not significant.
a 60Ge rotating source was used for attenuation correction. 15O-H2O PET studies were not adjusted for partial-volume effects because of increased variance associated with correction of relatively lower-resolution CBF images. A total of 238 CBF scans from 28 participants were available for analysis; 15 CBF scans were not available. The mean ± SD number of scans per participant was 8.5 ± 0.7 (range, 7–9). No significant differences between the proportions of unavailable studies for high- and low-11C-PIB-retention groups ($\chi^2; P = 1.0$) were observed.

**11C-PIB Studies**

Dynamic 11C-PIB PET studies were performed on average 10.8 ± 0.8 y after the first and 2.8 ± 0.8 y after the last CBF scan as part of the Baltimore Longitudinal Study of Aging neuroimaging study. Studies were performed on a GE Advance scanner in 3-dimensional mode, and 37 time frames (90 min) were obtained at rest. The PET scanning started immediately after an intravenous bolus injection of 529.1 ± 37 MBq (14.3 ± 1 mCi) of 11C-PIB with a specific activity of 152.8 ± 55.5 GBq/µmol (4.13 ± 1.5 Ci/µmol) (range, 36.26–249.01 GBq/µmol [0.98–6.73 Ci/µmol]). Participants were fitted with a thermoplastic mask for PET to minimize motion during scanning. Transmission scans in 2-dimensional mode were used for attenuation correction of the emission scans. Dynamic images were reconstructed using filtered backprojection with a ramp filter (image size, 128 × 128; pixel size, 2 × 2 mm; slice thickness, 4.25 mm), yielding a spatial resolution of about 4.50-mm full width at half maximum at the center of the field of view. We did not perform partial-volume correction on the 11C-PIB images, but the partial-volume effect was minimized by excluding edges of tissue in definition of volumes of interest. In view of the high noise level of voxel-based kinetics, the benefits of partial-volume correction of 11C-PIB images remain uncertain (17).

**MRI-Based Region-of-Interest (ROI) Definition**

Spoiled gradient-recalled MRI scans (124 slices; image matrix, 256 × 256; pixel size, 0.93 × 0.93 mm; slice thickness, 1.5 mm) were coregistered to the mean of the first 20-min dynamic PET images for each participant using the mutual information method in the Statistical Parametric Mapping software (SPM2; Wellcome Department of Cognitive Neurology). In addition to the cerebellum, which was used as a reference region, 15 ROIs (caudate, putamen, thalamus, lateral temporal, mesial temporal, orbital frontal, prefrontal, occipital, superior frontal, parietal, anterior cingulate, posterior cingulate, pons, midbrain, and white matter) were manually drawn on the coregistered MR images (18).

**Quantification of DVRs and Definition of High- and Low-11C-PIB-Retention Groups**

The DVRs of ROIs were estimated by simultaneous fitting of a reference-tissue model to the 15 measured ROI time–activity curves (19). In addition, DVR images were generated for voxel-wise analysis (17,19). The mean cortical DVR was calculated by averaging DVR values from orbitofrontal, prefrontal, superior frontal, parietal, lateral temporal, occipital, and anterior and posterior cingulate regions. The mean cortical DVR was used to form groups rather than used as a continuous variable because of the possibility that associations with rCBF change would not be uniform across the range of cortical DVR values. For example, no relationships may be evident in individuals with low 11C-PIB values. To separate individuals into 2 groups, we used the 2-group K-mean cluster analysis in SAS 9.1 (SAS Institute, Inc.).

**SPM Analysis of CBF Scans**

Using SPM2, PET CBF scans were realigned, spatially normalized into standard stereotactic space, and smoothed using a gaussian filter of 12, 12, and 12 mm in the x, y, and z planes, respectively. To control for variability in global flow, rCBF values at each voxel were ratio-adjusted to the mean global flow of 50 mL/100 g/min for each image. The image data were analyzed using voxel-by-voxel comparisons to assess differences in linear rCBF changes between the high- and low-11C-PIB groups over time across all data points. Group x time interactions of linear changes in rCBF were computed, covarying for sex and the time between the last rCBF study and the 11C-PIB study. Significant effects for each contrast were based on the magnitude ($P \leq 0.001$) and spatial extent (>200 mm$^3$) of differences between groups.

**Spatial Distribution of Differences in rCBF and 11C-PIB for High- Versus Low-11C-PIB Groups**

Using SPM2, 11C-PIB parametric DVR images were spatially normalized and smoothed using a gaussian filter of 8, 8, and 8 mm in the x, y, and z planes. A 2-group t test was used to assess differences between the high- and low-11C-PIB-retention groups. Significant effects were determined on the basis of magnitude ($P \leq 0.001$) and spatial extent (>200 mm$^3$). For graphical illustration, significant differences in longitudinal rCBF change between the high- and low-11C-PIB-retention groups were superimposed on r maps of differences in amyloid deposition between the high- and low-11C-PIB groups using the Amide software (version 3) (20). To determine whether regions showing significant group differences in rCBF change overlap with those showing differences in amyloid deposition, a restricted search analysis was performed limited to regions showing significant group differences in rCBF (mask threshold, $P = 0.05$).

**RESULTS**

**Study Participants**

High- and low-11C-PIB-retention groups were similar in age and other demographic characteristics and did not differ significantly in white matter signal abnormalities on MRI. Compared with the low-11C-PIB-retention group, the high-11C-PIB group showed a nonsignificant trend for lower scores on the Mini-Mental Status Examination (MMSE). In addition, mean CDR scores were higher in the high-11C-PIB-retention group (Table 1) than those in the low-11C-PIB group. At the time of the 11C-PIB study, the high-11C-PIB-retention group included only 1 individual (CDR = 0.5) who met consensus criteria for MCI. The remainder of the high-11C-PIB group included 4 individuals with a CDR of 0.5 who did not yet meet consensus criteria for MCI and 5 individuals with a CDR of 0. Four of the 5 individuals with a CDR of 0.5 received their initial rating of 0.5 during imaging visits before the 11C-PIB PET study, but none of the 4 progressed to a CDR greater than 0.5 during follow-up. During the 8 y of CBF imaging, none of the participants in either group developed central nervous system disease (stroke, epilepsy, brain tumor, Parkinson’s disease) or severe cardiac disease (myocardial infarction, coronary artery disease requiring angioplasty or bypass surgery).
Cluster Analysis of Mean Cortical DVR
To separate individuals into 2 groups, we used 2-group K-mean cluster analysis and used a cubic clustering criterion to test for uniformity within the data. This analysis identified 2 clusters separated at approximately 1.3 DVR, characterized by a cubic clustering criterion of 0.547, which indicates that they represent potential clusters (Fig. 1). On the basis of the cluster analysis, 10 participants were assigned to the high--11C-PIB-retention group and 18 participants were assigned to the low--11C-PIB-retention group. Significantly more participants in the high--11C-PIB-retention group than in the low--11C-PIB-retention group (P = 0.0126) had a CDR of 0.5 at the time of the 11C-PIB study (Table 1). Only 1 participant in the low--11C-PIB-retention group had a CDR score of 0.5, with a CDR sum of the boxes equal to 0.5. Individuals in the high--11C-PIB-retention group with a CDR of 0.5 had very mild cognitive impairment (mean CDR sum of boxes, 1.5 [±0.5]; maximum sum of boxes, 2).

Longitudinal Changes in High-- and Low--11C-PIB-Retention Groups
Longitudinal increases and decreases in rCBF differed significantly between high-- and low--11C-PIB-retention groups (P < 0.001; >200 mm³ spatial extent). Compared with the low--11C-PIB group, greater increases over time in rCBF in the high--11C-PIB group were seen in the left medial frontal (Brodmann’s area [BA] 10) and left inferior frontal gyrus (BA 45/46), right precuneus (BA 7), left inferior parietal lobule (BA 40), and left postcentral gyrus (BA 43). Greater decreases in rCBF over time in the high--11C-PIB group than in the low--11C-PIB group were found in the right anterior to middle cingulate (BA 32/24), right supramarginal gyrus (BA 40), left thalamus, and midbrain bilaterally (Table 2).

Voxel-based differences in amyloid deposition between the high-- and low--11C-PIB-retention groups are shown in Figure 2 in the lower row. The pattern of voxel-based differences in 11C-PIB retention between the groups is very similar to a map of 11C-PIB retention in a typical AD case. Superimposed on the voxel-based differences in 11C-PIB retention between the groups are regions of significant differences in longitudinal rCBF change between the groups. As confirmed by restricted search analysis, all regions, except brain stem and inferior thalamus, that show significant differences in longitudinal rCBF change between high-- and low--11C-PIB groups overlap regions of greater amyloid retention in the high--11C-PIB group.

DISCUSSION
This evaluation of older adults without dementia demonstrates regionally specific differences in longitudinal rCBF changes over time between individuals with and without amyloid deposition. In the years preceding measurement of amyloid deposition, both greater increases and greater decreases over time are observed in specific brain regions when individuals without dementia with high cortical 11C-PIB retention are compared with individuals with low 11C-PIB retention. The spatial distribution of rCBF changes in the high--11C-PIB group occurs primarily in areas with known amyloid deposition; however, not all areas that have been documented previously to have high 11C-PIB retention show longitudinal rCBF changes. Differences in longitudinal rCBF change between the high-- and low--11C-PIB-retention groups are seen in some structures that have altered rCBF in individuals with MCI who progress to AD (11,21). Unlike these cross-sectional imaging studies, the high--11C-PIB-retention group in our study is composed of a mix of healthy elderly and individuals with a CDR of 0.5, of whom only 1 has a consensus diagnosis of MCI. These differences in sample characteristics may account for differences between our findings and those of studies of individuals with MCI who progress to AD.

Greater declines in rCBF over time in the high-- than in the low--11C-PIB-retention group were found in the anterior cingulate gyrus extending to the right middle cingulate gyrus, the supramarginal gyrus, the thalamus, and the midbrain. Decreased rCBF has been previously reported

### TABLE 2

<table>
<thead>
<tr>
<th>Region</th>
<th>Brodmann’s area</th>
<th>Side</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>t value</th>
<th>P (uncorrected)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Greater linear increases in rCBF</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Precuneus</td>
<td>7</td>
<td>R</td>
<td>6</td>
<td>−48</td>
<td>44</td>
<td>3.55</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Inferior parietal lobule</td>
<td>40</td>
<td>L</td>
<td>−58</td>
<td>−22</td>
<td>28</td>
<td>3.25</td>
<td>0.001</td>
</tr>
<tr>
<td>Postcentral gyrus</td>
<td>43</td>
<td>L</td>
<td>−46</td>
<td>−12</td>
<td>22</td>
<td>3.58</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Medial frontal gyrus</td>
<td>10</td>
<td>L</td>
<td>−8</td>
<td>50</td>
<td>14</td>
<td>3.60</td>
<td>&lt;0.001</td>
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<tr>
<td>Inferior frontal gyrus</td>
<td>45/46</td>
<td>L</td>
<td>−32</td>
<td>28</td>
<td>12</td>
<td>4.41</td>
<td>&lt;0.001</td>
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<tr>
<td>Greater linear decreases in rCBF</td>
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<td></td>
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</tr>
<tr>
<td>Cingulate gyrus</td>
<td>32/24</td>
<td>R</td>
<td>16</td>
<td>22</td>
<td>28</td>
<td>4.51</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Supramarginal gyrus</td>
<td>40</td>
<td>R</td>
<td>48</td>
<td>−52</td>
<td>34</td>
<td>3.01</td>
<td>0.001</td>
</tr>
<tr>
<td>Thalamus</td>
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<td>−14</td>
<td>14</td>
<td>3.74</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>Midbrain</td>
<td>L</td>
<td>4</td>
<td>−24</td>
<td>−8</td>
<td>3.18</td>
<td>0.001</td>
<td></td>
</tr>
<tr>
<td>Midbrain</td>
<td>L</td>
<td>−6</td>
<td>−18</td>
<td>−2</td>
<td>3.12</td>
<td>0.001</td>
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</table>
in the anterior cingulate gyrus on SPECT of individuals who converted to AD (22), and deficits in executive function, in which the anterior cingulate has been reported to play a role, are among those exhibited early in the course of AD (23). In addition to the greater longitudinal decrease in the anterior cingulate for the high–11C-PIB-retention group, we also observed greater decreases over time for the high–11C-PIB group in the supramarginal gyrus and thalamus. CBF decreases in the supramarginal gyrus have been observed previously in cross-sectional studies comparing individuals with AD with controls (24,25) and in longitudinal follow-up of individuals with AD (25). SPECT studies have also shown early involvement of the thalamus in AD (22). We also found greater longitudinal declines in midbrain rCBF in the high–11C-PIB group, indicating greater decreases in neural activity in this region. We did not observe significant group differences in longitudinal rCBF changes for the posterior cingulate gyrus, an area showing early decreases in rCBF (22,25) and glucose metabolism (26) as well as increased amyloid deposition (27,28) in individuals diagnosed with MCI and AD.

Greater decreases in longitudinal rCBF in those with high amyloid deposition may represent the inability to further offset neuronal insult or neuronal death, a process that is also associated with decreased 18F-FDG activity in areas with high 11C-PIB deposition (1,28,29). In addition to focal decreases in neural activity, several other mechanisms either alone or in combination may explain decreases in CBF. These include compromise of the lumina of small vessels due to amyloid deposition itself or vasoconstrictive properties of endothelin-1, a molecule formed during interaction of Aβ with the receptor for advanced glycation end products (30); decreased synaptic integration (31); decreased choline acetyltransferase activity as an individual progresses toward AD (32); degeneration of indirect projections from the basal forebrain to cerebral microcirculation (33); and cell death due to amyloid toxicity (34).

We also found some regions that showed greater longitudinal increases in rCBF in those with higher amyloid load, compared with those with low amyloid load. These regions included inferior and medial frontal, postcentral, and inferior parietal lobule regions and the precuneus. All these regions with greater longitudinal rCBF increases also showed greater amyloid deposition in the high–11C-PIB-retention group in a voxel-based analysis. In one 11C-PIB PET study of individuals without dementia, the precuneus was reported to be the first region showing significant amyloid deposition (3). In addition to precuneus and posterior cingulate, the frontal regions have also shown pronounced 11C-PIB retention in prior studies (1,4–6,27,28). Notably, these neocortical regions also demonstrate early amyloid deposition in pathologic studies (7,8).

Although decline in neuronal function has been extensively documented, especially once an individual has been diagnosed with AD, accumulating evidence indicates potential increases in regional brain activity in the very early stages of cognitive impairment. Relative increases in rCBF
have been found in frontal regions in individuals who later progress to dementia, compared with controls and individuals with MCI who remain stable (21). Functional MRI studies of individuals with MCI also indicate initial increases in neuronal function early in the disease process (35). Moreover, neuropathologic studies of cognitively intact individuals whose postmortem findings are consistent with AD show hypertrophy of neuronal nuclei, nucleoli, and cell bodies in several brain regions, as compared with controls (36). These changes, which may be a marker of the resistance of the brain to the effects of amyloid deposition, would likely be associated with increased rCBF in view of increased metabolic demands. Although the concurrent relationship between rCBF and amyloid deposition has not yet, to our knowledge, been reported, our findings of greater longitudinal increase in rCBF in some brain regions of individuals with increased amyloid load is consistent with these other observations of increased neuronal activity in individuals with no or early MCI.

Although the specific mechanisms related to these longitudinal changes in rCBF remain unclear, either amyloid deposition itself or one of the pathophysiologic processes associated with amyloid deposition may play a role in the observed changes. One possible explanation for the pattern of greater longitudinal increases and decreases in rCBF in association with amyloid deposition may be that greater decreases in rCBF across time represent a response to neuronal insult that subsequently leads to neuronal death, whereas greater longitudinal increases in rCBF represent an attempt to preserve neuronal function despite an increase in amyloid load. These increases in rCBF, possibly representing neuronal plasticity changes, may play a role in the maintenance of cognitive function until such processes are overcome by increasing pathophysiology.

CONCLUSION

We demonstrate that the pattern of longitudinal rCBF changes in those with high 11C-PIB retention differs from those with low 11C-PIB retention. In this group of older adults without dementia, those with high 11C-PIB deposition show a decline in rCBF over the previous years, probably reflecting decrements in neuronal function. However, in the same group, there is also a greater increase in rCBF over time seen in other areas, suggesting that compensatory mechanisms aimed at preserving brain function are invoked in the setting of amyloid deposition. The spatial distribution of these changes corresponds to regions typically showing high amyloid deposition; however, not all areas of known amyloid deposition show changes in rCBF. Concurrent evaluation of longitudinal changes in rCBF and amyloid deposition with changes in cognitive status will be required to investigate in detail the interplay between CBF and amyloid deposition and their possible clinical implications. Further development, refinement, and application of different physiologic and molecular probes, in concert with longitudinal and perhaps more focused and refined clinical observations, will bring us closer to understanding the pathophysiology of MCI and AD and may aid in determining which individuals at risk for AD are most likely to progress to disease.

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