Spatial patterns of brain amyloid-β burden and atrophy rate associations in mild cognitive impairment

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Amyloid-β accumulation in the brain is thought to be one of the earliest events in Alzheimer’s disease, possibly leading to synaptic dysfunction, neurodegeneration and cognitive/functional decline. The earliest detectable changes seen with neuroimaging appear to be amyloid-β accumulation detected by 11C-labelled Pittsburgh compound B positron emission tomography imaging. However, some individuals tolerate high brain amyloid-β loads without developing symptoms, while others progressively decline, suggesting that events in the brain downstream from amyloid-β deposition, such as regional brain atrophy rates, play an important role. The main purpose of this study was to understand the relationship between the regional distributions of increased amyloid-β and the regional distribution of increased brain atrophy rates in patients with mild cognitive impairment. To simultaneously capture the spatial distributions of amyloid-β and brain atrophy rates, we employed the statistical concept of parallel independent component analysis, an effective method for joint analysis of multimodal imaging data. Parallel independent component analysis identified significant relationships between two patterns of amyloid-β deposition and atrophy rates: (i) increased amyloid-β burden in the left precuneus/cuneus and medial-temporal regions was associated with increased brain atrophy rates in the left medial-temporal and parietal regions; and (ii) in contrast, increased amyloid-β burden in bilateral precuneus/cuneus and parietal regions was associated with increased brain atrophy rates in the right medial temporal regions. The spatial distribution of increased amyloid-β and the associated spatial distribution of increased brain atrophy rates embrace a characteristic pattern of brain structures known for a high vulnerability to Alzheimer’s disease pathology, encouraging for the use of 11C-labelled Pittsburgh compound B positron emission tomography measures as early indicators of Alzheimer’s disease.
These results may begin to shed light on the mechanisms by which amyloid-β deposition leads to neurodegeneration and cognitive decline and the development of a more specific Alzheimer’s disease-specific imaging signature for diagnosis and use of this knowledge in the development of new anti-therapies for Alzheimer’s disease.

**Keywords:** MRI; $^{11}$C-PiB PET; Alzheimer’s disease; mild cognitive impairment; amyloid-β; amyloid; brain atrophy rate; multimodal brain imaging  
**Abbreviations:** ADNI = Alzheimer’s Disease Neuroimaging Initiative; $^{11}$C-PiB = $^{11}$C-labelled Pittsburgh compound B

## Introduction

The formation of amyloid plaques in the brain, mainly consisting of insoluble amyloid-β protein fragments, is thought to be a major factor that leads to degradation of neurons and ultimately to the development of Alzheimer’s disease (Braak and Braak, 1991). Amyloid-β plaques, together with neurofibrillary tangles, are also the hallmarks of a definite diagnosis of Alzheimer’s disease at autopsy (Braak and Braak, 1991). The development of radiotracers, such as the $^{11}$C-labelled Pittsburgh compound B ($^{11}$C-PiB), which binds to amyloid-β plaques, has made it possible to study plaque accumulation in life using $^{11}$C-PiB PET (Klunk et al., 2004; Mintun, 2005; Price et al., 2005; Lockhart, 2006; Nordberg, 2007, 2008). Several $^{11}$C-PiB PET studies have demonstrated increased brain amyloid-β deposition in patients with Alzheimer’s disease (Price et al., 2005; Kemppainen et al., 2006; Edison et al., 2007; Mikho et al., 2008; Frisoni et al., 2009; Tolboom et al., 2009b; Devanand et al., 2010). A growing number of $^{11}$C-PiB PET studies have also reported increased brain amyloid-β in subjects with mild cognitive impairment (Price et al., 2005; Kemppainen et al., 2007; Pike et al., 2007; Rowe et al., 2007; Forsberg et al., 2008; Tolboom et al., 2009b), individuals who fall between normal ageing and dementia and who have an increased risk of developing Alzheimer’s disease (Petersen et al., 2009). Furthermore, studies of cognitively normal elderly subjects suggest that the presence of amyloid-β, as detected by $^{11}$C-PiB PET, is associated with a risk of developing symptomatic Alzheimer’s disease (Klunk et al., 2004; Mintun et al., 2006; Kemppainen et al., 2007; Morris et al., 2009). These results have raised the possibility that $^{11}$C-PiB PET may detect preclinical Alzheimer’s disease when treatment intervention may be most effective. However, about 10–30% of cognitively normal elderly subjects have amyloid-β positive findings on $^{11}$C-PiB PET images even without apparent cognitive deficits (Galvin et al., 2005; Driscoll et al., 2006; Mintun et al., 2006; Pike et al., 2007; Rowe et al., 2007; Aizenstein et al., 2008; Villemagne et al., 2008). Whether cognitively normal subjects with high levels of $^{11}$C-PiB binding will develop Alzheimer’s disease later on is unknown. Furthermore, only about 50% of patients with mild cognitive impairment advance to Alzheimer’s disease within a short period of time (Lopez et al., 2007) although 60–65% of all patients diagnosed with mild cognitive impairment are PiB-positive. The observations that some individuals tolerate high brain amyloid-β loads without cognitive deficits, while others suffer cognitive decline and there is a wide range in the rate of decline, suggest amyloid-β plaques alone are not sufficient to cause cognitive impairment and accompanying neurodegeneration. In this study, we aim to shed light on the role of amyloid-β in neurodegeneration by elucidating the relationship between regional amyloid-β retention and progression of regional brain atrophy in subjects with clinically established cognitive impairments. We chose subjects diagnosed with mild cognitive impairment to test whether or not such a relationship can be detected in mild stages of brain atrophy. Furthermore, to detect both spatially localized as well as spatially distributed effects of amyloid-β retention on the progression of regional brain atrophy we utilized novel multivariate statistical methods for image analysis.

Over the past few years, several imaging studies have assessed amyloid-β deposition and brain atrophy together across a wide range of cognitive impairments using MRI and $^{11}$C-PiB PET (Archer et al., 2006; Fotenos et al., 2008; Jack et al., 2008b; Mormino et al., 2009; Bourgeat et al., 2010; Chetelat et al., 2010). While a study of a healthy elderly population reported no associations between amyloid-β burden and regional rate of brain volume decline in the preceding years (Driscoll et al., 2010), other studies have found an association between amyloid-β levels and brain tissue atrophy and ventricular expansion at mild stages of cognitive deficits (Jack et al., 2009; Chetelat et al., 2010). Even in stages of advanced cognitive impairment (e.g. after conversion to Alzheimer’s disease), discrepant findings regarding an association between amyloid-β levels and brain atrophy have been reported (Archer et al., 2006; Chetelat et al., 2010). Though several factors might have contributed to these mixed findings, two aspects that are crucial for a better understanding of amyloid-β and brain atrophy relations were not addressed. First, no previous imaging study has determined to what extent the increased regional amyloid-β aggregation relates to increased rates of regional brain atrophy. Second, and perhaps more importantly, no previous study has investigated to what extent the regional variations in amyloid-β aggregation relate to regional variations in increased rates of brain atrophy across brain structures.

The main purpose of this study was to understand the relationship between the regional distribution of amyloid-β burden and the regional pattern of brain atrophy rates in mild cognitive impairment. Specifically, we tested two hypotheses: (i) increased amyloid-β levels in mild cognitive impairment are associated with increased brain atrophy rates; and (ii) the spatial distribution of elevated amyloid-β in mild cognitive impairment is associated with a characteristic pattern of increased and spatially distributed atrophy rates across brain structures typically involved in Alzheimer’s disease pathology, such as the medial temporal lobe,
precuneus and posterior cingulate cortices. Conventional image statistics based on region of interests or voxel-wise tests, such as statistical parametric mapping (Worsley et al., 2004) are generally limited to studies of localized relationships, i.e. when amyloid-β deposition leads to brain atrophy at the same location, but are inadequate to capture spatially distributed relations across the brain. To overcome this limitation, we turned to the concept of parallel independent component analysis, an effective method for joint analysis of multimodality imaging data (Calhoun and Adali, 2009). Parallel independent component analysis identifies not only variations across both image modalities but also variations across brain regions by evaluating all regional distributions simultaneously, thereby taking intrinsic relationships between these distributions into consideration. Therefore, our findings should lead to deeper insight into the biology of the disease than methods that evaluate the modalities as well as each brain region separately.

Materials and methods

Participants

The 61 participants diagnosed with mild cognitive impairment in this study were recruited between 2005 and 2008 through the Alzheimer’s Disease Neuroimaging Initiative (ADNI) from 56 centres in the USA and Canada (Table 1) (Petersen et al., 2010). The ADNI was funded as a prospective, longitudinal study to identify biomarkers of early Alzheimer’s disease for clinical trials, supported by the National Institute on Ageing (NIA), the National Institute of Biomedical Imaging and Bioengineering (NIBIB), the Food and Drug Administration (FDA), private pharmaceutical companies and non-profit organizations, as a 5-year public-private partnership. Written consent was obtained from all subjects participating in the study, and the study was approved by the institutional review board at each participating site. This study also complied with the Health Insurance Portability and Accountability Act guidelines.

The subjects with mild cognitive impairment were over the age of 55 years with memory complaints and memory difficulties that were verified by an informant, scored below the education adjusted cut-off on a baseline Logical Memory II subscale (Delayed Paragraph Recall) from the Wechsler Memory Scale—Revised (Wechsler, 1987), scored between 24 and 30 on a baseline Mini-Mental State Exam (Folstein et al., 1975), scored a 0.5 on the Clinical Dementia Rating scale (Morris, 1993), scored at least 0.5 on the Memory Box and scored ≤12 on the 17-item Hamilton Depression rating scale at the ADNI study screening. General cognition and functional performance of each participant was sufficiently preserved such that a diagnosis of Alzheimer’s disease could not be made by the site physician at the time of the screening visit. Participants with any significant neurological disease other than suspected incipient Alzheimer’s disease, such as Parkinson’s disease, multi-infarct dementia, Huntington’s disease, normal pressure hydrocephalus, brain tumour, progressive supra-nuclear palsy, seizure disorder, subdural haematoma, multiple sclerosis or history of significant head trauma followed by persistent neurological defaults or known structural brain abnormalities as well as subjects with baseline MRI scans with evidence of infection, infarction, or other focal lesions, and with multiple lacunes or lacunes in a critical memory structure were excluded from the study. Further details regarding inclusion and exclusion criteria can be found at www.adni-info.org.

Table 1 Study group demographics

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<tbody>
<tr>
<td>Age at 11C-PiB PET imaging baseline, years</td>
<td>75.5 (8.0)</td>
</tr>
<tr>
<td>Follow-up, years</td>
<td>1.0 (0.1)</td>
</tr>
<tr>
<td>Gender, % female</td>
<td>34</td>
</tr>
<tr>
<td>Education, years</td>
<td>16.4 (2.7)</td>
</tr>
<tr>
<td>MMSE at 11C-PiB PET imaging baseline</td>
<td>27.0 (2.4)</td>
</tr>
<tr>
<td>CD Scale at 11C-PiB PET imaging baseline</td>
<td>1.8 (2.1)</td>
</tr>
<tr>
<td>CDR at 11C-PiB PET imaging baseline</td>
<td>0.5 (0.2)</td>
</tr>
<tr>
<td>ADAS-Cog at 11C-PiB PET imaging baseline</td>
<td>11.2 (5.6)</td>
</tr>
<tr>
<td>Percent of APOE e4 carriers</td>
<td>61</td>
</tr>
</tbody>
</table>

Statistics are provided in mean (standard deviation) format. ADAS-Cog = Alzheimer Disease Assessment Scale-Cognitive; CDR = Clinical Dementia Rating; GD Scale = Geriatric Depression Scale; MMSE = Mini-Mental State Exam.
Magnetic resonance imaging acquisition

The participants at each site underwent the following standardized 1.5 T MRI protocol: two $T_1$-weighted MRI scans, using a sagittal volumetric magnetization prepared rapid gradient echo (MP-RAGE) sequence, with an echo time of 4 ms, repetition time of 9 ms, flip angle of 8° and acquisition matrix size of $256 \times 256 \times 166$ in the x-, y- and z-dimensions with a nominal voxel size of $0.94 \times 0.94 \times 1.2$ mm. A designated centre selected the MP-RAGE image with higher quality and corrected for system-specific image artefacts such as geometry distortion, B1 non-uniformity and intensity inhomogeneities (Jack et al., 2008a). A k-space post-processing technique called ‘auto-correction’ was implemented to correct 3D image fields that have been corrupted by bulk head motion (McGee et al., 1997). The technique operates by iteratively adding phase offsets to select lines in k-space and then automatically evaluating a quantitative metric in image space such as entropy or gradient entropy. The presence, absence and severity of common artefacts (e.g. blurring due to head motion) are also indicated in the MRI quality control files. MRI scan at the time of the baseline $^{11}$C-PiB PET scanning was considered as the baseline structural scan in this study and the follow-up structural scan within 1 year of this baseline structural scan was used to quantify the annual brain tissue loss rate as described below.

Unbiased magnetic resonance imaging structural brain image template

To avoid bias towards a particular subject’s geometry in generating a reference brain image, we used the baseline MRI data from all subjects in this study to create an unbiased large deformation structural brain image template (Joshi et al., 2004). First, the skull, scalp, extra-cranial tissue, cerebellum and brain stem (at the level of the diencephalon) were removed from each $T_1$-weighted image data using the automated Brain Surface Extraction software (Shattuck and Leahy, 2002), followed by quality check. The automatically generated Brain Surface Extraction brain masks were manually refined if necessary. A study-specific unbiased large deformation template was then created using baseline structural brain images from all 61 participants with mild cognitive impairment by applying a fluid registration algorithm as described in full elsewhere (Joshi et al., 2004; Lorenzen et al., 2005). Briefly, unbiased large deformation template formation incorporates an unbiased approach where all baseline brain images are first simultaneously affine (i.e. 12 degrees-of-freedom) transformed to adjust for global variations in brain positioning and scale. Affine transformed subject brain images are then simultaneously deformed iteratively to yield an average unbiased large deformation template. A non-linear inverse-consistent fluid-flow deformation field from each affine transformed subject brain image to the resulting unbiased large deformation template is estimated through this process.

Creation of maps of brain atrophy rates

To compute the rate of regional brain volume change (i.e. brain atrophy rate), the 1-year follow-up $T_1$-weighted MRI scan of each subject was first brain tissue masked using Brain Surface Extraction and then the data were rigid body transformed into the image space of the corresponding baseline scan. An inverse-consistent diffeomorphic fluid-flow warping algorithm was further applied to spatially normalize the registered 1-year follow-up image to the baseline brain image. Finally, rates of regional brain volume change were computed for each subject based on the determinant of the Jacobians describing the diffeomorphic deformation fields (i.e. Jac-map), yielding maps of rates of fractional volume contraction or expansion at each voxel in each subject’s native baseline image space.

$^{11}$C-labelled Pittsburgh compound B positron emission tomography and magnetic resonance imaging data fusion

For the joint analysis of maps of PiB-standardized uptake value ratio and brain atrophy rates, the baseline PiB-standardized uptake value ratio map (co-registered to the corresponding baseline structural magnetic resonance image) and the corresponding Jac-map of each subject were transformed to the unbiased large deformation template space using the same affine and unbiased large-deformation fluid-flow field estimated in creation of the unbiased large deformation brain template as described above and as illustrated in Fig. 1. In addition, the PiB-standardized uptake value ratio map and Jac-map were each smoothed with a Gaussian spatial kernel of 4 and 6 mm full-width at half-maximum, respectively. The 2:3 ratio of the different smoothing kernels for PiB-standardized uptake value ratio map and Jac-map was estimated according to methods described in Hagler et al. (2006) to achieve comparable degrees of smoothing for the two image modalities.

Joint statistical analysis of $^{11}$C-labelled Pittsburgh compound B positron emission tomography and magnetic resonance imaging

In this study, we aimed to test distributed relationships between amyloid-β accumulation and 1-year brain atrophy rates across brain regions. To achieve this goal, we analysed both image modalities jointly and furthermore took all image voxels into account simultaneously using the concept of parallel independent component analysis. This analytical strategy is designed to test the hypothesis that a systematic pattern of regional amyloid-β accumulation, as reflected by PiB-standardized uptake value ratio maps, is associated with a regional pattern of 1-year brain atrophy rates, as measured with MRI Jac-maps. Importantly, this analysis makes no restrictions on local relationships between PiB-standardized uptake value ratio and Jac-map but also includes relationships that might be spatially dissociated, involving influence of amyloid-β accumulation on brain atrophy rates in distant brain regions.

The mathematical foundations of parallel independent component analysis are described in detail in Liu et al. (2009). We used the parallel independent component analysis package FIT (Fusion ICA Toolbox http://icatb.sourceforge.net) to obtain statistical maps of joint relationships. In short, parallel independent component analysis applied to multimodality imaging data aims to identify independent components in each image modality as well as the relationships of these independent components across image modalities.

In the context of this study, the parallel independent component analysis design identified spatially independent components in each PiB-standardized uptake value ratio and Jac-map of the MRI while simultaneously revealing the largest variations across the subjects with mild cognitive impairment that PiB-standardized uptake value ratio and MRI Jac-map had in common. The number of significant
independent components in each modality was estimated using both
the Akaike information criterion and the minimum description length
criterion (Calhoun et al., 2001). For each modality, the loading par-
parameters expressing the contribution of each independent component
to the variance across subjects were estimated. Each independent
component for each modality was scaled to unit standard deviation,
yielding z-score maps in the unbiased large deformation template brain
image space. All component maps were thresholded at a z-score level
of $|z| > 2.5$ (99.4% cumulative probability) for visualization purposes.

Based on these loading parameters, we computed Pearson’s correl-
ation coefficients for all pairs of PiB-standardized uptake value ratio
and brain atrophy rates independent components while further ac-
counting for variations in age, gender, education levels, APOE ε4 car-
rier status, and Alzheimer Disease Assessment Scale-Cognitive test
scores across the subjects with mild cognitive impairment. Alzheimer
disease Assessment Scale-Cognitive test scores were included in the
model to adjust for different levels of cognitive impairment within the
mild cognitive impairment group, which might skew the regional pat-
terns of amyloid-β accumulation and brain atrophy rates as well as the
relationships between them. The Pearson’s correlation coefficients
were then used to identify significant relationships between regional
brain amyloid-β accumulation and brain atrophy rates after correction
for multiple comparison using a false discovery rate at significance
level of $q < 0.05$ (Benjamini and Hochberg, 1995).

**Results**

We found two significant and spatially distributed component
pairs in mild cognitive impairment, each depicting an association
between PiB-standardized uptake value ratio measures at baseline
and regional brain atrophy rates over 1 year follow-up.

The spatial extent of the first significant component pair is
shown in Fig. 2, overlaid onto the unbiased large deformation brain
template. The first component pair showed a partial correl-
ation of 0.77 between baseline PiB-standardized uptake value ratio
and regional brain atrophy rates with a false discovery rate cor-
rected significance level of $P = 0.0004$. The adjusted $R^2$ value of
the fitted model for this component pair was 0.58 with $P < 10^{-4}$.
The brain regions identified by this first component pair, as shown
in Table 2, imply more amyloid-β burden was associated with
higher rates of brain atrophy. Regions with increased amyloid-β
burden included the precuneus, cuneus, medial temporal
(including fusiform, lingual, entorhinal), inferior parietal regions
in the left hemisphere, bilateral posterior cingulate and right lateral
fronto-orbital regions. Interestingly, a predominantly left-sided in-
crease in amyloid-β burden in these brain regions was associated
with higher atrophy rates (i.e. more grey matter and white matter
tissue loss and larger sulcal cerebro-spinal fluid expansion over a
1-year period) in the medial temporal regions (including entorh-
inal, fusiform, parahippocampal), amygdala, inferior temporal, su-
perior frontal, inferior frontal in the left hemisphere, bilateral
posterior cingulate and left cerebellum.

We further found a second significant component pair. The
spatial extent of the second significant component pair is shown
in Fig. 3, overlaid onto the unbiased large deformation brain
template. The adjusted $R^2$ value of the fitted model for the second
component pair was 0.34 ($P < 0.0001$) and the partial correlation
coefficient between the baseline PiB-standardized uptake value ratio
and regional brain atrophy rates was 0.65 (false discovery rate corrected $P = 0.04$). In the second component pair, increased
amyloid-β burden in bilateral precuneus/cuneus, bilateral posterior
cingulate, left inferior frontal, left inferior parietal, left superior parietal and bilateral thalamus regions was associated with higher atrophy rates in the right medial temporal regions (including entorhinal, fusiform, parahippocampal, subcallosal), right amygdala, right middle fronto-orbital, right gyrus rectus and left parahippocampus, as shown in Table 2. Note, each of the two significant component pairs from the parallel independent component analysis identifies a spatially distributed pattern of regions with high amyloid-β burden and regions with high rates of brain atrophy in mild cognitive impairment, as seen in the voxel-wise overlay in Figs 2 and 3.

We also explored the possibility that the relationship between PiB-standardized uptake value ratio measures and brain atrophy rates were not simply an artefact of partial-volume effects of PiB data. We tested this by computing a grey matter index from tissue segmented MRI data obtained at baseline for each region of interest where a significant relationship between PiB-standardized uptake value ratio and brain atrophy rates were found and then repeating the analysis with the grey matter index as an additional covariate to account for partial-volume effects of PiB data. We found that the inclusion of a grey matter index reduced relations by ~1% (where significance of variability due to partial-volume

<table>
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<tr>
<th>Table 2 Regions of the associated baseline amyloid-β and 1-year brain atrophy rates components identified by parallel independent component analysis</th>
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<tbody>
<tr>
<td><strong>Baseline PiB-standardized uptake value ratio</strong></td>
</tr>
<tr>
<td>Region</td>
</tr>
<tr>
<td>First parallel independent component analysis component pair</td>
</tr>
<tr>
<td>Left precuneus/cuneus</td>
</tr>
<tr>
<td>Left inferior parietal</td>
</tr>
<tr>
<td>Bilateral posterior cingulate</td>
</tr>
<tr>
<td>Right lateral fronto-orbital</td>
</tr>
<tr>
<td>Left medial temporal (fusiform, lingual, entorhinal)</td>
</tr>
<tr>
<td>Second parallel independent component analysis component pair</td>
</tr>
<tr>
<td>Bilateral precuneus/cuneus</td>
</tr>
<tr>
<td>Bilateral posterior cingulate</td>
</tr>
<tr>
<td>Left inferior frontal</td>
</tr>
<tr>
<td>Left inferior parietal</td>
</tr>
<tr>
<td>Left superior parietal</td>
</tr>
<tr>
<td>Bilateral thalamus regions</td>
</tr>
</tbody>
</table>

Figure 2  First parallel independent component analysis component pair. SUVR = standardized uptake value ratio.
effect was \( P = 0.60 \) for the first component of parallel independent components analysis and by 7% (where significance of variability due to partial-volume effect was \( P = 0.17 \)) for the second component, suggesting that the relationships between amyloid-\( \beta \) load and brain atrophy rates cannot simply be explained as partial-volume artefacts.

**Discussion**

There are two major findings in this study: first, high amyloid-\( \beta \) levels, detected by \( ^{11} \text{C-PiB} \) PET imaging, in mild cognitive impairment are associated with increased brain atrophy rates, detected by longitudinal MRI. The finding adds to the generally held view that patients with mild cognitive impairment who have amyloid-\( \beta \) pathology have high brain atrophy rates and are likely to progress to Alzheimer’s disease within a relatively short time, while subjects with mild cognitive impairment without amyloid-\( \beta \) pathology have much lower brain atrophy rates and may not develop Alzheimer’s disease. Second, the spatial distribution of increased amyloid-\( \beta \) and the associated spatial distribution of increased brain atrophy rates embrace a characteristic pattern of brain structures known for a high vulnerability to Alzheimer’s disease pathology.

The finding of an association between increased amyloid-\( \beta \) and increased brain atrophy rates in mild cognitive impairment implies that aggregated amyloid-\( \beta \) deposition leads to a progressive loss of brain tissue over time as opposed to a static loss from random insults. Although a progressive tissue loss due to increased amyloid-\( \beta \) deposition is expected (Jack et al., 2009), an interesting observation of our study is that the regional distribution of increased amyloid-\( \beta \) and that of increased rates of brain tissue loss are not necessarily overlapping in mild cognitive impairment. Specifically, the joint analysis of \( ^{11} \text{C-PiB} \) PET and structural MRI data identified two component pairs of spatially distributed relations between amyloid-\( \beta \) deposition and brain atrophy rates. In the first component pair, high amyloid-\( \beta \) burden in the left precuneus/cuneus and posterior cingulate regions was associated with increased rates of brain tissue loss primarily in the left medial temporal lobe, including the entorhinal cortex and parahippocampal gyrus. On the other hand, in the second component pair, high amyloid-\( \beta \) burden in bilateral precuneus/cuneus and posterior cingulate region was associated with increased rates of brain tissue loss primarily in the right medial temporal lobe regions. Taken together, these results may begin to shed light on the mechanisms by which regionally selective amyloid-\( \beta \) deposition in mild cognitive impairment leads to a spatial pattern of distributed neurodegeneration, which resembles that seen in Alzheimer’s disease. Potentially, these mechanisms are anterograde and selectively target the memory networks. The systematic patterns of increased amyloid-\( \beta \) burden and increased brain atrophy rates in the patients with mild cognitive impairment, strongly resembling the pattern typically seen in Alzheimer’s disease, further supports the view that these relations reflect brain alterations presymptomatic to Alzheimer’s disease and could be useful for staging disease severity and for monitoring treatment interventions in preclinical stages of Alzheimer’s disease.

Structural MRI studies in Alzheimer’s disease have consistently revealed a pattern of neuroanatomical abnormalities that predominately involve structures in the medial temporal cortex (i.e. hippocampus and the entorhinal cortex) (Du et al., 2001, 2003, 2004; deToledo-Morrell et al., 2004; Thompson et al., 2004; Hampel et al., 2005; Stoub et al., 2005; Morra et al., 2008, 2009a, b; Schroeter et al., 2009) where the early pathological changes are seen, then gradually extends to temporoparietal cortical areas (Chetelat and Baron, 2003; Whitwell et al., 2007, 2008; Desikan et al., 2008; Hua et al., 2008) as severity of Alzheimer’s disease progresses (Jack et al., 2004, 2005, 2008; deCarli et al., 2007; Whitwell et al., 2007, 2008). Findings in both parallel independent component analysis component pairs point to a selective vulnerability of these regions to Alzheimer’s disease pathology, consistent with histopathological findings. The finding in mild cognitive impairment that a localized distribution of amyloid-\( \beta \) deposition is...
associated with a characteristic pattern of brain atrophy that resembles the atrophy pattern seen in Alzheimer’s disease is encouraging for the use of $^{11}$C-PiB-PET measures as early indicators of Alzheimer’s disease. Most importantly, elucidating the detrimental relationship between the local amyloid-β burden and rates of brain atrophy is of great interest to enhance our understanding of the underlying mechanisms of disease and use of this knowledge in development of new anti-therapies for Alzheimer’s disease.

Another important observation is the prominence of the left hemisphere in the relationship between increased amyloid-β burden and increased rates of brain atrophy. This observation is consistent with previous reports of higher atrophy rates on the left hemisphere compared with the right hemisphere in patients with Alzheimer’s disease (Thompson et al., 2003). It is also interesting in comparison to other MRI studies, which found a trend of higher right than left asymmetry of hippocampal atrophy rates in cognitively normal elderly subjects and some evidence that suggests there is a change in asymmetry during the progression toward Alzheimer’s disease (Barnes et al., 2005). It is therefore possible that the laterality in brain atrophy rates that we observed in the first parallel independent component analysis component pair is another potentially useful index for staging Alzheimer’s disease severity as well as for assessing disease-altering interventions. Further prospective studies of joint $^{11}$C-PiB PET and MRI evaluations including elderly individuals without cognitive deficits are warranted to determine the value of asymmetric findings for an early detection of Alzheimer’s disease pathology.

Another interesting observation in this study is a marked dissimilarity between the spatial distributions of increased amyloid-β deposition and that of increased brain atrophy rates in mild cognitive impairment. In particular, the spatial dissimilarity between increased amyloid-β deposition that included the precuneus and cingulate cortex whereas increased atrophy rates involved mainly medial temporal lobe structures but left precuneus and the cingulate cortex largely unaffected, is unexpected. It is possible that the brain regions with increased amyloid-β deposition in patients with mild cognitive impairment has already reached an equilibrium or plateau in terms of the rate of atrophy very early in the course of Alzheimer’s disease while atrophy rates in regions with low amyloid-β burden are further progressing. This interpretation is also consistent with the paradigm that amyloid-β is the first indication of disease and higher atrophy rates are a downstream process. From a functional anatomy point of view, the precuneus is implicated in the recollection of past episodes whereas posterior cingulate projecting from thalamus and neocortex to entorhinal cortex via cingulum fibres is characteristically active during recall and deactivated during encoding of episodic memory (Cabeza and Nyberg, 2000). Cognitive decline in episodic memory is one of the earliest clinical syndromes of Alzheimer’s disease. A popular hypothesis on disease mechanism suggests that amyloid-β accumulation is responsible for brain atrophy and hence the cognitive decline (Jack et al., 2010). According to this hypothesis, our findings together with the functional role of precuneus and posterior cingulate regions suggest that anti-amyloid-β therapy might be successful when administered very early in the disease evolution before the synaptic and neuronal loss in brain regions susceptible to early amyloid-β deposition reach a plateau. In a related investigation, Driscoll et al. (2010) reported no association between amyloid-β burden and rates of brain atrophy in the preceding years in healthy elderly individuals. Their finding suggests that either ageing-related structural brain atrophy rates are independent of amyloid-β deposition, or amyloid-β deposition must reach a particular level and be present for an extended time period before causing extensive Alzheimer’s disease-related brain atrophy. To further elucidate these hypotheses future studies focusing on especially early mild cognitive impairment populations are needed. In addition, the use of a selective in vivo marker of regional aggregated tau deposition in the form of neurofibrillary tangles would be useful to assess the linkage between regional amyloid-β, regional neurofibrillary tangle deposition and regional atrophy rates. To date, no useful neurofibrillary tangle-specific imaging agent has been conclusively demonstrated.

An important conceptual difference between our study and other imaging studies in mild cognitive impairment is the approach to jointly evaluate not only variations across the two imaging modalities, but also variations that the imaging modalities have in common across brain regions. This approach provided insight into the spatial distributions of amyloid-β and brain atrophy rate relations that conventional analysis methods, which test relationships by regions of interest or voxel-by-voxel, cannot provide. The difference in methods may also explain the differences in findings between our studies and others. For example, other $^{11}$C-PiB-PET studies found higher amyloid-β depositions also in other brain regions, including the frontal, posterior cingulate, parietal, and lateral temporal cortices, as well as in the subcortical regions including caudate and putamen (Kemppainen et al., 2007; Rowe et al., 2007). Similarly, other MRI studies in mild cognitive impairment have also found significant atrophy in temporal lobe white matter, orbitofrontal and medial prefrontal grey matter, posterior cingulate, insula and uncus (Apostolova et al., 2006; Becker et al., 2006; Colliot et al., 2008; Fan et al., 2008). However, the variations in amyloid-β and brain atrophy rates in these regions might be independent of each other, which would explain our inability to detect significant variations in these regions with parallel independent component analysis. Although another study reported no significant relationship between increased amyloid-β and brain atrophy in mild cognitive impairment, the authors suggested a concordance between atrophy and amyloid-β deposition in the posterior cingulate, precuneus and lateral temporoparietal association cortices during early stages of the disease process (Chetelat et al., 2010). Lack of an amyloid-β-brain atrophy rate association, especially in posterior cingulate and precuneus regions, is a surprising finding of our study. An explanation would be that reduced $^{11}$C-PiB signal due to atrophy in these regions would affect our ability to detect a local relationship between amyloid-β deposition and atrophy progression, since we chose to analyse non-atrophy-corrected PiB-standardized uptake value ratio data.

Amyloid-β levels in CSF have also been proposed as a diagnostic biomarker for Alzheimer’s disease by various researchers (Andreasen et al., 1999; Blennow and Hampel, 2003; Clark et al., 2003; Bouwman et al., 2007; De Meyer et al., 2010; Fjell et al., 2010). Recently, we reported that lower CSF amyloid-β$_{42}$ concentrations were associated with higher rates of brain volume.
loss in the left temporal and parietal cortices (Tosun et al., 2010) in mild cognitive impairment, similar to our findings of higher amyloid-β levels in the brain from this study. Furthermore, significant correlation ($r = -0.72$) between the global $^{11}$C-PiB binding load and the CSF amyloid-β$_{1-42}$ concentrations were found across healthy elderly, mild cognitive impairment and Alzheimer's disease populations (Jagust et al., 2009; Tolboom et al., 2009a). Taken together, these findings also encourage the use of $^{11}$C-PiB-PET imaging measures as early indicators of Alzheimer's disease.

In Alzheimer's disease, the default-mode network is markedly disconnected compared with healthy elderly controls, especially in the bilateral calcarine/cuneus, bilateral precuneus/post-cingulate, left lingual, left middle temporal gyrus, left parahippocampal, right angular, right dorsolateral prefrontal cortices as well as in the left hippocampus (Greicius et al., 2004; Zhou et al., 2010). It has been hypothesized that disruption of functional connectivity within default-mode network could be an alternative biomarker to structural alterations for early diagnosis of Alzheimer's disease (Hedden et al., 2009; Yvette et al., 2010). Moreover, it has been suggested that destabilization of neuronal networks and compensatory responses contribute to cognitive impairments in Alzheimer’s disease and that such destabilization stems, at least in part, from aberrant increases in neuronal activity induced by high amyloid-β levels (Palop and Mucke, 2009). Regions with high amyloid-β deposition and associated accelerated atrophy also mirror the regions affected in default-mode network. In particular, posterior components of the default-mode network, including the precuneus and posterior cingulate, are particularly vulnerable to early deposition of amyloid-β (Braak and Braak, 1991) and the temporal components of the default-mode network are mentioned in regional atrophy early in the disease process. One interpretation of these observations is that amyloid-β pathology could be linked to neural dysfunction in a distributed network of brain regions linked to memory function (Sperling et al., 2009). An important question is if default-mode network dysfunction is a functional consequence of amyloid-β deposition, especially since increased neuronal activity has been shown to increase the production, release and/or accumulation of amyloid-β, raising the possibility of a vicious cycle in which amyloid-β promotes its own production through alterations in neuronal network activity (Kamenetz et al., 2003; Cirrito et al., 2005). Further longitudinal studies are required to better understand such a causality between amyloid-β deposition and default-mode network dysfunction.

Several limitations of our study ought to be mentioned. First, mild cognitive impairment subjects are notoriously a very heterogeneous group and other pathologies, unrelated to Alzheimer’s disease, may have contributed to variations in both amyloid-β and rates of brain atrophy. In addition, we did not separate patients with mild cognitive impairment with high PiB burden and high atrophy rates from those with low PiB burden and low atrophy rates to maintain sufficient statistical power. By analysing the groups together, we may have underestimated the extent of joint variations in amyloid-β and atrophy rate in the high PiB burden/atrophy rate group of mild cognitive impairment and similarly, overestimated these variations in the low PiB burden/atrophy rate group. The value of our findings to predict Alzheimer’s disease remains uncertain because the number of patients with mild cognitive impairment who had $^{11}$C-PiB PET and converted to Alzheimer’s disease was small at the time of this study. In addition, the number of cognitively normal subjects in ADNI, who had both $^{11}$C-PiB PET and sequential MRI scans at the time of the study was too small for a reliable statistical analysis. Therefore, the extent to which our findings separate mild cognitive impairment from cognitively normal subjects or have value to predict who will develop mild cognitive impairment is unclear.

Our study also has several technical limitations. First, errors in image registrations between the image modalities as well as errors in spatial normalization may have increased variability and hence diminished statistical power to detect intrinsic relationships between atrophy rates and amyloid-β burden at a localized level. Second, the current framework of parallel independent component analysis assumes that measurements in each image voxel are independent and noise is identically distributed, which is likely not true and therefore statistical evaluations could be inflated.

In conclusion, our major finding links a greater amyloid-β deposition in the precuneus, a region generally known for early accumulation of amyloid plaques, to a unique pattern of increased brain atrophy within each hemisphere, resembling the pattern seen also in Alzheimer’s disease. These results may begin to shed light on the mechanisms by which amyloid-β deposition leads to neurodegeneration and cognitive decline and the development of a more specific Alzheimer’s disease-specific imaging signature for diagnosis.

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