Voxel-Based Morphometric Comparison Between Early- and Late-Onset Mild Alzheimer's Disease and Assessment of Diagnostic Performance of Z Score Images

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BACKGROUND AND PURPOSE: Voxel-based morphometry (VBM), used for detecting brain atrophy, permits comparison of local gray matter concentration at every voxel in an image between two groups. We sought to delineate the specific patterns of cerebral gray matter loss with regard to onset of Alzheimer’s disease (AD) by using MR imaging and VBM and to evaluate the diagnostic performance of VBM with Z score images.

METHODS: Two groups of 30 patients with mild AD of different ages of onset were examined. Mean ages in the early- and late-onset groups were 60.2 ± 5.2 and 71.5 ± 2.6 years, respectively. Control subjects were aged-matched healthy volunteers. Regions of gray matter loss in early- and late-onset AD were examined with VBM. Diagnostic performance of Z score images obtained with the VBM method was evaluated in patients and control subjects by calculating the area under the receiver operating characteristic curve ($A_z$).

RESULTS: Both AD groups had significantly reduced gray matter in the bilateral medial temporal regions. In addition, the early-onset group had more severe gray matter loss in the bilateral parietal and posterior cingulate cortices and precuneus region. No difference was noted in diagnostic performance of Z score images between the early- ($A_z = 0.9435$) and late-onset ($A_z = 0.9018$) groups.

CONCLUSION: Differences were noted in the patterns of regional gray matter loss in patients with early-onset AD versus those with late-onset AD. Parietotemporal and posterior cingulate gray matter loss was found in early-onset AD but not in late-onset AD. Z score images obtained with VBM had a great diagnostic performance for mild AD and can be applied for detecting mild AD in clinical examinations.

The classification of Alzheimer’s disease (AD) into subtypes based on the time of onset of symptoms is still controversial (1, 2). Clinically, patients with early-onset AD have greater degrees of other cognitive dysfunction and a rapid progression of cognitive deficits rather than memory disturbance in the early stage of the disease, whereas those with late-onset AD have a greater degree of memory disturbance. Histopathologic studies demonstrate the different pathologic changes in these two subtypes (3), and 2-[$\text{fluorine}$-$18$] fluoro-2-deoxy-D-glucose positron emission tomographic studies show metabolic differences between the two subtypes, indicating that the parietotemporal and posterior cingulate hypometabolic dysfunction in the early-onset type is much greater than that in the late-onset type (4–8). In structural imaging studies, such as CT and MR imaging, structural differences between the two subtypes have not been well investigated. This may be because it is difficult to measure and investigate regional atrophy in the brain by using only MR imaging or CT. Recently, voxel-based morphometry (VBM) of T1-weighted MR images has been used to investigate gray matter concentration in some diseases. VBM is a new technique for detecting brain atrophy that permits the comparison of local gray matter concentration at every voxel in an image between two groups of subjects (9). Some authors have reported on AD...
abnormalities by using this method (10, 11); however, they did not compare early-onset with late-onset AD. We investigated whether cerebral gray matter loss is different between early- and late-onset AD by using the VBM method with MR images and assessed the clinical diagnostic performance of the VBM method for mild AD.

**Methods**

**Subjects**

Thirty subjects in each AD subgroup (early or late onset) composed the statistical analysis set and a prototypic template set. Twenty additional subjects in each AD subgroup composed a receiver operating characteristic (ROC) test set. Before the examination, written informed consent was obtained from all the patients and/or their relatives and from all the volunteer subjects. The study protocol was approved by our institution's Ethical Committee.

**Patients.** We selected two groups of 30 patients with mild AD in whom the onset of dementia was before (early onset) or after (late onset) the age of 65 years, according to the criteria of the National Institute of Neurologic and Communicative Disorders and Stroke/Alzheimer's Disease and Related Disorders Association (NINCDS/ADRDA) (12) (Table 1, first group). The Mini-Mental State Examination (MMSE) scores (13) of the patients with early-onset AD and those with late-onset AD were matched.

In addition, another two groups consisting of 20 patients with early-onset AD and 20 patients with late-onset AD were recruited for diagnostic evaluation with use of the VBM method (Table 1, second group). Both patient groups fulfilled the criteria of the NINCDS/ADRDA for probable AD (12).

All patients underwent examination by neuropsychologists and psychiatrists, MR imaging, MR angiography of the neck and head, electroencephalography, laboratory tests, and neuropsychologic tests. To determine the age at onset, family members, caregivers, and friends were asked when they first noted any mental or behavioral changes in the patients.

**Healthy control subjects.** Two groups of healthy age-matched volunteer subjects were selected for each group of patients with AD (Table 1, first group). Also, another two groups of healthy volunteers were selected for the two subtypes of AD for diagnostic evaluation with use of the VBM method (Table 1, second group). The control subjects showed no clinical evidence of cognitive deficits or neurologic disease and were not taking short- or long-term drug therapy at the time of the imaging examinations. They had no abnormal findings on MR images, disregarding age-related atrophy and white matter change on T2-weighted images.

**MR Imaging Procedures**

The MR imager was a 1.5-T Signa Horizon (GE Medical Systems, Milwaukee, WI). Sagittal, coronal, and axial T1-weighted spin-echo (SE) images (550/15/2 [TR/TE/NEX], 5-mm thickness, 2.5-mm gap) and axial T2-weighted fast SE images (3000/21,105/2) were obtained for diagnosis. Then, coronal 3D spoiled gradient-echo (SPGR) imaging (14/5/2, 20° flip angle, 220-mm field of view, 256 × 256 matrix, 124 × 1.5-mm contiguous sections) was performed for VBM analysis.

**Data Analysis**

Statistical parametric mapping (SPM) 99 software (Wellcome Department of Cognitive Neurology, London, United Kingdom) was installed on a Windows computer. Calculations and image matrix manipulations were performed in MATLAB 5.3 (MathWorks Inc., Natick, MA). For VBM analysis with SPM 99, all the coronal SPGR MR imaging data sets were reconstructed to axial data sets and then converted to ANALYZE format and displayed with the right hemisphere on the right.

Anatomic normalization and statistical processing were performed with SPM 99 software. All the individual MR images were transformed into a standard stereotactic anatomic space, and then the images were automatically segmented by using a cluster analysis technique (9). This process partitioned the images into gray matter, white matter, and CSF by using a modified mixture model cluster analysis technique with a correction for image intensity nonuniformity. The image sets were smoothed with an isotropic Gaussian filter (12-mm full-width at half-maximum), and individual global gray matter densities were normalized by proportional scaling.

**Correlation between aging and gray matter loss in healthy subjects.** First, we examined the correlation between the regional gray matter density and aging in the control group by using the SPM statistical technique.

**Regional gray matter loss in the early- versus late-onset AD groups.** Next, the regional gray matter densities in the early-onset group versus that in the younger control group and in the late-onset group versus that in the older control group were compared. Then, for comparing the early-onset, EO, and late-onset, LO, groups directly, we used the following SPM contrast, which means a set of subjects for comparison and a statistical design in the SPM statistics, for group comparison to remove the aging effect (8):

\[
(EO \text{ younger control}) - (LO \text{ older control}), (\text{EO} - \text{young control}) - (\text{LO} - \text{older control}), (\text{EO} - \text{young control}) - (\text{LO} - \text{older control}), (\text{EO} - \text{young control}) - (\text{LO} - \text{older control}), (\text{EO} - \text{young control}) - (\text{LO} - \text{older control})
\]

Significance was accepted if the voxels survived a Bonferroni-corrected threshold of \( P < .05 \) or an uncorrected threshold of \( P < .001 \).

**Diagnostic value in the early- versus late-onset AD groups.** For evaluation of diagnostic utility, Z score map inspections were performed (14). The following process was performed with free software, Easy Z Score Imaging System (eZIS; Daiichi Radiotisotope Laboratory, Tokyo, Japan). Each prototypic early-onset AD and late-onset AD template map was obtained from the first comparison between each AD subtype and the age-matched control subjects in the first group in this study, which indicated the area where the gray matter density was significantly reduced in each early-onset AD and late-onset AD map at the threshold of \( P = .001 \). Two normal databases were constructed by averaging, on a voxel-by-voxel basis, the image sets of each of the first 30 healthy subjects for the younger and older control groups.

Next, Z scores were calculated for each voxel of each subject in the second group by using the two normal databases: Z score = \(((\text{normal mean}) - (\text{individual value})) / (\text{normal SD}))\). The subset of pixels exceeding a threshold of \( Z > 2.0 \) for each individual was displayed on the prototypic template and used as a Z score map image. Then, Z score map images of the subjects in the second group were classified as follows: no pixels over Z

**TABLE 1: Characteristics of patients with AD and healthy control subjects**

<table>
<thead>
<tr>
<th>Group</th>
<th>Sex (F:M)</th>
<th>Age (y)*</th>
<th>MMSE Score*</th>
</tr>
</thead>
<tbody>
<tr>
<td>First</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early-onset AD</td>
<td>22:8</td>
<td>60.2 ± 5.2</td>
<td>23.0 ± 2.1</td>
</tr>
<tr>
<td>Younger controls</td>
<td>20:10</td>
<td>59.6 ± 3.8</td>
<td>29.9 ± 0.3</td>
</tr>
<tr>
<td>Late-onset AD</td>
<td>22:8</td>
<td>71.5 ± 2.6</td>
<td>22.3 ± 1.8</td>
</tr>
<tr>
<td>Older controls</td>
<td>20:10</td>
<td>71.4 ± 3.5</td>
<td>29.4 ± 0.9</td>
</tr>
<tr>
<td>Second</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early-onset AD</td>
<td>14:6</td>
<td>60.8 ± 4.6</td>
<td>23.5 ± 1.9</td>
</tr>
<tr>
<td>Younger controls</td>
<td>17:3</td>
<td>59.1 ± 2.7</td>
<td>29.8 ± 0.8</td>
</tr>
<tr>
<td>Late-onset AD</td>
<td>17:3</td>
<td>72.2 ± 3.2</td>
<td>23.4 ± 2.0</td>
</tr>
<tr>
<td>Older controls</td>
<td>11:9</td>
<td>70.3 ± 4.2</td>
<td>29.5 ± 0.8</td>
</tr>
</tbody>
</table>

* Data are mean ± SD.
Correlation Between Aging and Gray Matter Loss in Healthy Subjects

Gray matter densities in the bilateral perisylvian cortices, parahippocampal gyri, and pre- and postcentral gyri were significantly and negatively correlated with age in the healthy control group \((P < .05, \text{corrected})\). No positive correlation was noted between gray matter density and age.

Regional Gray Matter Loss in Early- versus Late-Onset AD Groups

Figure 2 shows the specific voxels that had significantly more gray matter loss in each group of patients with AD compared with the age-matched control subjects \((P < .001, \text{uncorrected})\). Regions of significant atrophy seen in both groups were located in the bilateral medial temporal lobes. The differences between the late-onset group and its age-matched control subjects were not as great as the differences between the early-onset group and its age-matched control subjects. In addition, in the early-onset group, the bilateral inferior parietal lobules, precuneus, perisylvian cortices, and basal forebrain region, and the right inferior frontal gyrus gray matter densities were significantly decreased compared with those of the younger control group (Table 3).

Between the two AD groups, the gray matter densities in the bilateral precuneus, left parietal cortices, right middle temporal gyrus, and left fusiform gyrus were lower in the early-onset group than in the late-onset group (Fig 3, Table 4). The late-onset group did not show significantly decreased gray matter density compared with that in the early-onset group at the threshold of \(P < .001\), uncorrected.

Diagnostic Value in the Early- versus Late-Onset Groups

Figure 4 shows the ROC curves and SE of the Z score images in a 54-year-old patient with early-onset AD (MMSE score = 23). Mild parietal lobular atrophy can be detected by visual inspection of the conventional T1-weighted image (Fig 5A) compared with that of a 57-year-old healthy control subject (MMSE score = 30) (Fig 6A). By using the Z score map, it is easy to detect the region and degree of atrophy in this early-onset AD case (Fig 5B) and to see that there is no significant region in the healthy subject (Fig 6B).

Figure 7 shows conventional MR images and Z score images in a 73-year-old healthy control subject (MMSE score = 30) shows no hippocampal atrophy but the left parietal lobe seems to be atro-

TABLE 2: Location of greatest gray matter reduction in healthy control subjects in relation to age

<table>
<thead>
<tr>
<th>Location</th>
<th>Z Value</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Right perisylvian cortices</td>
<td>7.12</td>
<td>40</td>
<td>13</td>
<td>4</td>
</tr>
<tr>
<td>Right hypothalamus</td>
<td>6.19</td>
<td>3</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>Left perisylvian cortices</td>
<td>6.06</td>
<td>34</td>
<td>19</td>
<td>1</td>
</tr>
<tr>
<td>Left parahippocampal gyrus</td>
<td>5.77</td>
<td>19</td>
<td>44</td>
<td>4</td>
</tr>
<tr>
<td>Left postcentral gyrus</td>
<td>5.47</td>
<td>47</td>
<td>23</td>
<td>41</td>
</tr>
<tr>
<td>Right postcentral gyrus</td>
<td>5.41</td>
<td>36</td>
<td>36</td>
<td>49</td>
</tr>
</tbody>
</table>

score 2 on the significant area of the prototypic template: definite normal, 1; clusters consisting of more than 2 pixels \((4 \times 2 \text{ mm})\) of maximum Z score 2.0–2.5 existed on the significant area of prototypic template: probable normal, 2; Z score 2.6–3.0: indeterminate, 3; Z score 3.1–4.0: probable AD, 4; and Z score over 4.1: definite AD, 5.

We performed ROC analysis, and the area under the ROC curve \((A_r)\) values and their standard errors (SE) were calculated by using ROCKIT software 0.9B (Metz CE, Department of Radiology, University of Chicago, IL).

Results

Correlation Between Aging and Gray Matter Loss in Healthy Subjects

Gray matter densities in the bilateral perisylvian cortices, parahippocampal gyri, and pre- and postcentral gyri were significantly and negatively correlated with age in the healthy control group \((P < .05, \text{corrected})\) (Fig 1, Table 2). No positive correlation was noted between gray matter density and age.

Regional Gray Matter Loss in Early- versus Late-Onset AD Groups

Figure 2 shows the specific voxels that had significantly more gray matter loss in each group of patients
phied, whereas the Z score map shows no significant atrophy in the parietal lobe (Fig 8B).

**Discussion**

In the present study with use of the MR imaging VBM method, we demonstrated differences in patterns of regional gray matter loss in patients with early- versus those with late-onset AD and the possibility of clinical application of Z score images obtained by using the MR imaging VBM method for diagnosis of AD.

Many studies incorporating structural images like MR images have been used to evaluate neurodegenerative changes in AD (15). Most of them used volumetry of the hippocampal formation, parahippocampal gyri, and entorhinal cortices. However, this kind of study requires detailed tracing of the target struc-

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**TABLE 3: Location of the greatest gray matter reduction in the cluster regions of significant gray matter density reduction in early-onset and late-onset AD groups compared with age-, sex-, and severity-matched control subjects**

<table>
<thead>
<tr>
<th>Group and Location</th>
<th>Z Value</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td>YHC &gt; EO</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right parahippocampal gyrus</td>
<td>5.79</td>
<td>17</td>
<td>-36</td>
<td>0</td>
</tr>
<tr>
<td>Left inferior parietal lobe</td>
<td>5.06</td>
<td>-45</td>
<td>-50</td>
<td>41</td>
</tr>
<tr>
<td>Right inferior parietal lobe</td>
<td>4.77</td>
<td>38</td>
<td>-62</td>
<td>41</td>
</tr>
<tr>
<td>Left operculum</td>
<td>4.72</td>
<td>-41</td>
<td>-13</td>
<td>-6</td>
</tr>
<tr>
<td>Right inferior frontal gyrus</td>
<td>4.50</td>
<td>41</td>
<td>8</td>
<td>30</td>
</tr>
<tr>
<td>Right precuneus</td>
<td>4.25</td>
<td>15</td>
<td>-65</td>
<td>14</td>
</tr>
<tr>
<td>Left inferior parietal lobe</td>
<td>3.96</td>
<td>-47</td>
<td>-58</td>
<td>23</td>
</tr>
<tr>
<td>OHC &gt; LO</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Left hippocampus</td>
<td>5.13</td>
<td>-26</td>
<td>-7</td>
<td>-16</td>
</tr>
<tr>
<td>Right hippocampus</td>
<td>4.35</td>
<td>25</td>
<td>-38</td>
<td>-4</td>
</tr>
</tbody>
</table>

Note.—YHC indicates younger healthy controls; EO, early-onset AD; OHC, older healthy controls; LO, late-onset AD. Threshold is \( P < .001 \), uncorrected.

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**Fig 2.** A. Statistical parametric maps show comparison of patients with early-onset AD with age-matched healthy volunteers (the younger control subjects). Highlighted areas are regions of significant gray matter loss in the patients with early-onset AD compared with age-matched control subjects at a threshold of \( P < .001 \), uncorrected. Bilateral medial temporal lobes, inferior parietal lobules, precuneus, and perisylvian cortices and the right inferior frontal gyrus and bilateral cingulate cortex are highlighted. L indicates left; R, right.

B. Statistical parametric maps show comparison of patients with late-onset AD and age-matched healthy volunteers (the older control subjects). Highlighted areas are regions of significant gray matter loss in patients with late-onset AD compared with age-matched control subjects at a threshold of \( P < .001 \), uncorrected. Bilateral medial temporal cortices are highlighted. L indicates left; R, right.
than that for late-onset AD (10). Investigated gray matter loss in age-related loss of the substantia innominata (17).

To minimize the effects of age, sex, and cognitive impairment in our study, the patients with AD were divided into two subtypes (early and late onset), with the age and sex well matched between the patients and control subjects, and the cognitive impairment matched between the early- and late-onset AD groups. As previous functional imaging studies have demonstrated, patients with AD show a typical pattern of hypometabolism bilaterally in the parietotemporal cortices, posterior cingulate cortices, and sometimes frontal association cortices, associated with relatively preserved metabolism and blood flow in the visual cortices, the sensorimotor cortices, cerebellum, and subcortical structures (18–20). Small et al (21) and Sakamoto et al (8) found that metabolism in the parietal lobe was lower in their early-onset group than in their late-onset group. Morphologic evaluation in our study also demonstrated that parietal regions were not significantly affected in the gray matter in the late-onset group compared with the age-matched control group. It is well known that the clinical symptoms of patients with early- and those with late-onset AD are different (22). Patients with late-onset AD are more severely impaired in memory and in naming items than are those with early-onset AD. Neuronal loss in the CA zone of the Ammon horn and in the subiculum is greater in patients with late-onset AD than in those with early-onset AD (23). Although healthy control subjects lose 12% of their hippocampal pyramidal neurons between ages 50 and 70 years, patients with AD lose an estimated 57% of their neurons over the same time period (24). The severity of memory impairment in the late-onset subgroup may be related to the relatively severe pathologic changes in these regions. However, our study did not demonstrate significantly greater medial temporal gray matter loss in the late-onset group than in the early-onset group, because the magnitude of medial temporal cortical loss in the late-onset AD group was not significant. Although, when comparing with the age-matched control subjects, the magnitude and expansion of significant gray matter loss was larger in the late-onset group versus the control group than in the early-onset group versus the control group. There was laterality to some of our findings: the right inferior frontal gyrus in early-onset group versus the control subjects; and the left parietal cortex, right angular gyrus, and left fusiform gyrus in the early-onset group versus the late-onset group. Some asymmetry of cortical atrophy and metabolic or perfusional reduction in AD is not rare (25, 26).

The application of VBM for examination of the atrophied brain is still controversial (27, 28). Recently, however, VBM was applied in many studies of AD and related diseases, and reasonable findings
were obtained (10, 11, 16, 29–33). In the current study, a structural image combined with a statistical Z score image was first applied to obtain a clinical diagnosis of mild AD. For practical clinical examination, the VBM method requires a relatively short time for obtaining a final statistical map, the Z score image. A combination of a statistical map like our Z score image with the VBM method provides an accurate diagnosis in patients with AD. Diagnosis based on the annual change derived from serial MR images may be superior to the performance of a diagnosis based on a single MR image, although our study demonstrated a good result by using a single MR image.

As our study showed that there were differences in regional cortical gray matter concentration reductions between the early- and late-onset groups, it is better to set up more than two normal age-referenced databases for this kind of VBM method when making a clinical diagnosis of AD. Our structural analysis enabled accurate detection of mild AD, indicating that the VBM method coupled with Z score images can be applied as a clinical tool for diagnosing AD.

In almost all imaging studies of dementia, including this study, imaging findings are compared with clinically known cases of AD. In such a study, it is impossible for the imaging study to be more sensitive or specific than the clinical examination. A way to over-
come this is by following up the patients over a number of years and establishing at the start of the study the imaging parameters that characterize those patients who go on to develop AD, as demonstrated by De Leon et al (34). This clearly presented study provides a powerful demonstration of the ability of modern image analysis and MR images to characterize dementia and to help diagnose AD. This method of study design should be tested in assessing diagnostic performance with VBM and Z score images in future studies.

**Conclusion**

We performed a voxel-based analysis of regional gray matter loss in patients with early- versus those with late-onset AD. Our study results support previous pathologic evidence and show that there are differences in the alteration of regional gray matter loss between early- and late-onset AD, suggesting that AD should be classified into these two subtypes. The VBM method with Z score map can be applied for detecting mild AD in routine MR imaging examinations.

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