Tissue Microstructural Changes Are Independently Associated With Cognitive Impairment in Cerebral Amyloid Angiopathy

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Background and Purpose—Cerebral amyloid angiopathy (CAA) is a major cause of lobar intracerebral hemorrhage and cognitive impairment and is associated with white matter hyperintensities and cerebral microbleeds. MRI diffusion tensor imaging detects microstructural tissue damage in advanced CAA even in areas that appear normal on conventional MRI. We hypothesized that higher global mean apparent diffusion coefficient (mean ADC), reflecting a higher amount of chronic tissue disruption caused by CAA, would be independently associated with CAA-related cognitive impairment.

Methods—Preintracerebral hemorrhage cognitive impairment was systematically assessed using a standardized questionnaire (IQCODE) in 49 patients. Volume of white matter hyperintensities, number of microbleeds, and mean ADC were determined from MRIs obtained within 14.0–22.5 days of intracerebral hemorrhage cognitive impairment. White matter hyperintensities and mean ADC were measured in the hemisphere uninvolved by intracerebral hemorrhage to avoid confounding.

Results—Preintracerebral hemorrhage cognitive impairment was identified in 10 of 49 subjects. Mean ADC was the only variable associated with preintracerebral hemorrhage cognitive impairment and was elevated in those with preintracerebral hemorrhage cognitive impairment compared with those without (12.4±10−4 versus 11.7±10−4 mm²/s; P=0.03). Mean ADC positively correlated with age but not white matter hyperintensities or number of microbleeds. In logistic regression controlling for age and visible cerebral atrophy, mean ADC was independently associated with preintracerebral hemorrhage cognitive impairment (OR per 1×10−4 mm²/s increase=2.45, 95% CI 1.11 to 5.40, P=0.04).

Conclusions—Mean ADC is independently associated with preintracerebral hemorrhage cognitive impairment in CAA. The lack of correlation with other MRI markers of CAA suggests that mean ADC may be sensitive to distinct aspects of CAA pathology and its tissue consequences. These results suggest that global MRI diffusion changes are sensitive to clinically relevant microstructural alterations and may be useful markers of CAA-related tissue damage. (Stroke. 2008;39:1988-1992.)

Key Words: cerebral amyloid angiopathy • cerebral microhemorrhage • cognitive impairment • dementia • diffusion • microbleeds • white matter disease

Cerebral amyloid angiopathy (CAA) is a major cause of lobar intracerebral hemorrhage (ICH) and cognitive impairment in the elderly and is associated with a high prevalence of markers of small vessel disease, including white matter hyperintensities (WMH) and cerebral microbleeds (MB).1–4

ICH in sporadic CAA is likely caused by vessel fragility and rupture due to the deposition of amyloid within the media and adventitia of small- to medium-sized cerebral arteries.5 The presence of WMH visualized on conventional MRI sequences4 suggests that vascular amyloid deposition may alter white matter perfusion by causing stenosis or vascular dysfunction.6 This hypothesis is supported by the observations of extensive white matter damage in families with rare hereditary forms of CAA due to point mutations within the amyloid precursor protein.7,8 WMH have been associated with cognitive impairment in sporadic CAA.4

MRI diffusion tensor imaging has been used in other cerebral small vessel diseases to detect microstructural changes in cerebral tissue, even in areas that appear normal on conventional MRI,9–11 and to detect significant differences...
between CAA and control populations. Global quantitative measures can be obtained with diffusion tensor imaging by using whole-brain histograms of mean diffusivity such as mean apparent diffusion coefficient (ADC). Two recent publications, by independent research groups, showed that whole-brain ADC histograms were closely correlated with clinical disability in the rare hereditary small vessel disease cerebral autosomal-dominant arteriopathy with subcortical ischemic leukoencephalopathy and cerebral autosomal-dominant arteriopathy with subcortical ischemic leukoencephalopathy, is also a well-defined cerebral small vessel disease with a high prevalence of subcortical tissue lesions. We hypothesized that a similar relationship might be present between these global diffusion tensor imaging measures and CAA-related clinical disability. We therefore sought to determine whether higher global mean ADC would be associated with chronic cognitive impairment in CAA, independent of the effects of hemorrhagic stroke, possibly by reflecting a higher amount of chronic tissue disruption caused by the disease.

Methods

Subjects

The study subjects were recruited from an ongoing single-center prospective longitudinal cohort study of CAA. Subjects were consecutive patients aged ≥55 years who presented to the Massachusetts General Hospital between August 2001 and August 2005 with symptomatic lobar ICH and had MRI with diffusion-weighted sequences within 90 days of the index event. MRI was obtained at the discretion of the treating physician and was performed according to departmental protocol. Those with other potential causes of hemorrhage such as cavernous angioma or other vascular malformations, or hemorrhages in locations more typical of hypertensive ICH such as the basal ganglia and thalamus, were excluded. CAA was diagnosed according to the previously validated Boston Criteria; there were 5 subjects (10%) with definite CAA by tissue diagnosis, 19 subjects (39%) with probable CAA (2 or more ICH, including microbleeds on MRI, restricted to lobar regions), and 25 subjects (51%) with possible CAA (a single lobar ICH without MRI evidence of other hemorrhages).

Baseline clinical and demographic information was collected blinded to imaging data. Hypertension was defined as previous diagnosis of hypertension (>140/90 mm Hg) or use of antihypertensive treatment for control of blood pressure. Like in prior studies, we measured cognitive impairment before ICH to avoid the major influence on cognition introduced by occurrence of the hemorrhagic stroke itself. Pre-ICH cognitive impairment (PICI) was defined as the presence of deficits in memory or other cognitive areas sufficient to interfere with tasks of daily living before ICH and was assessed based on interview with the subject and informants, review of the medical records, and results of a standardized questionnaire (IQCODE short form) that included items related to memory, praxis, calculation, and reasoning. The IQCODE short form is a microangiopathy that affects cerebral small vessels diffusely and symmetrically in the brain, measurement of mean ADC values in one hemisphere should be reflective of whole-brain tissue microstructural change. Briefly, histograms of ADC values from ADC maps were generated for each patient using a bin width equal to 0.1×10⁻⁶ mm²/s. Voxels containing cerebrospinal fluid or areas of hemorrhage were excluded in all patients before calculation using a previously established superior threshold value of 27×10⁻⁶ mm²/s. To correct for cross-subject differences in brain volume, each histogram was normalized to the total number of hemispheric tissue voxels. The mean ADC derived from each histogram was used for analysis.

The volume of WMH was determined on fluid-attenuated inversion recovery sequence according to our previously published method. Using MRIcro software (University of Nottingham School of Psychology, Nottingham, UK; www.mricro.com), a region-of-interest map of supratentorial WMH in the hemisphere contralateral to the hematoma was created by signal intensity thresholding followed by manual editing as necessary. To correct for head size, we used the sagittal midline cross-sectional intracranial area as a surrogate measure of the intracranial volume. WMH volume was normalized to head size (nWMH) by dividing the subject’s WMH volume by the ratio of the subject’s intracranial area to the mean intracranial area of the study population. The number of hemorrhages, including MB, was determined on the gradient-echo sequence. A global visible cerebral atrophy score was calculated on T2- and fluid-attenuated inversion recovery sequences by summing individual scores (range, 0 to 3) in 5 separate brain regions to give a total score of 0 to 15. ICH volume was determined by segmentation of the ICH on the baseline CT scan, at the time of hemorrhagic stroke, using Alice software (Parexel Corporation, Waltham, Mass). We have previously shown high interrater reliability, by intraclass correlation coefficient (ICC), for measurements of nWMH (ICC = 0.98), midsagittal intracranial area (ICC = 0.92), MRI hemorrhages (ICC = 0.97), visible cerebral atrophy score (ICC = 0.79), and ICH volume (ICC = 0.99).

All imaging analyses were performed by raters blinded to clinical information.

Statistical Analysis

Mean ADC was compared with the presence or absence of PICI by Student t-test. The numbers of MRI microhemorrhages were divided into 3 categories (0, 1 to 4, or ≥5) to divide groups into tertiles and tested for association with PICI by Fisher exact test. Because the distribution of nWMH and MRI hemorrhages were right-skewed, Spearman’s correlations were used for comparisons with other continuous variables and Wilcoxon’s rank sum tests were used for comparisons with dichotomous variables.

Multivariate logistic regression models were used to find predictors of PICI, candidate covariates were those associated with PICI in univariate analysis (P<0.10). The final model variables were selected by the stepwise regression analysis with entry and removal with the guidelines of the Institutional Review Boards of Massachusetts General Hospital.

Image Acquisition and Analysis

MR images were acquired on 1.5-Tesla Sigma scanners (GE Medical Systems, Milwaukee, Wis) as previously described. Diffusion-weighted images (TR/TE 7500/97 to 99 ms, slice thickness 5 to 6 mm, interslice gap 1.0 mm, 128×128; b value=1000 s/mm²; 23 slices) were performed. Diffusion-weighted imaging scans were acquired in orthogonal directions and then averaged to make ADC measurements largely independent of the effects of anisotropic diffusion. Fluid-attenuated inversion recovery images (TR/TE 10 000/140 ms, slice thickness 5 mm, interslice gap 1.0 mm, 256×256) and axial gradient-echo images were obtained as previously described. MRIs were performed an average of 14.0 days after ICH (range, 0 to 89 days).

Mean ADC was determined in the hemisphere contralateral to the hematoma using a previously described method. Because CAA is a microangiopathy that affects cerebral small vessels diffusely and symmetrically in the brain, measurement of mean ADC values in one hemisphere should be reflective of whole-brain tissue microstructural change. Briefly, histograms of ADC values from ADC maps were generated for each patient using a bin width equal to 0.1×10⁻⁶ mm²/s. Voxels containing cerebrospinal fluid or areas of hemorrhage were excluded in all patients before calculation using a previously established superior threshold value of 27×10⁻⁶ mm²/s. To correct for cross-subject differences in brain volume, each histogram was normalized to the total number of hemispheric tissue voxels. The mean ADC derived from each histogram was used for analysis.
Visible cerebral atrophy, but this did not reach statistical significance (Table).

Mean ADC was not associated with nWMH ($P=0.44$) or the number of MB ($P=0.38$). The presence or absence of a history of hypertension was not significantly associated with mean ADC ($P=0.13$). We found no association between ICH volume and mean ADC values in the non-ICH hemisphere. The side of the hematoma did not have a significant effect on mean ADC values in the non-ICH hemisphere. Patients with higher mean ADC values were older ($r=0.51$, $P<0.001$) and had a higher visible cerebral atrophy score ($r=0.48$, $P<0.001$).

Clinical variables and MRI measures that were correlated with PICI in univariate analyses ($P<0.10$) were entered into a stepwise multivariate logistic regression model to determine independent predictors of PICI in CAA. In the final maximally adjusted model controlling for age and amount of visible cerebral atrophy, only mean ADC was independently associated with PICI (OR per $1\times10^{-4}$ mm$^2$/s increase = 2.45, 95% CI 1.11 to 5.40, $P=0.04$). The effect of visible cerebral atrophy on PICI was not significant in the final model ($P=0.33$). Other MRI markers such as nWMH and number of MB also did not have an independent effect on PICI. Other clinical variables, including history of hypertension and coronary artery disease, were not associated with PICI in multivariate models. Excluding patients with a history of ICH ($n=3$) from these analyses did not significantly change our results. In all of these analyses, there was no scanner effect that modified the interpretation of the results (data not shown).

**Discussion**

The major finding from this cohort study of patients with possible or probable CAA is that, among MRI markers of CAA, global mean ADC (reflecting microstructural tissue organization) is most strongly related to the presence of pre-ICH cognitive impairment. The association of mean ADC with PICI was independent of age, clinical variables, amount of visible cerebral atrophy, and other MRI markers. By contrast, we failed to detect a relationship between PICI and WMH volume, number of MB, or visible cerebral atrophy.

Decreased cognitive performance has previously been associated with CAA. The prevalence of cognitive dysfunc-

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**Table. Baseline Characteristics of Subjects in the Cohort by the Presence or Absence of PICI**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>PICI</th>
<th>No PICI</th>
<th>$P$ Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>6 (60)</td>
<td>21 (54)</td>
<td>0.99</td>
</tr>
<tr>
<td>Female</td>
<td>4 (40)</td>
<td>18 (46)</td>
<td></td>
</tr>
<tr>
<td>Age, years ± SD</td>
<td>75.8±6.4</td>
<td>73.5±8.3</td>
<td>0.44</td>
</tr>
<tr>
<td>History of hypertension</td>
<td>8 (80.0)</td>
<td>25 (64.1)</td>
<td>0.46</td>
</tr>
<tr>
<td>Atrial fibrillation</td>
<td>1 (10.0)</td>
<td>7 (18.0)</td>
<td>0.99</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>3 (30.0)</td>
<td>3 (7.7)</td>
<td></td>
</tr>
<tr>
<td>History of coronary artery disease</td>
<td>2 (20.0)</td>
<td>4 (10.3)</td>
<td>0.59</td>
</tr>
<tr>
<td>History of ischemic stroke</td>
<td>1 (10.0)</td>
<td>1 (2.6)</td>
<td>0.37</td>
</tr>
<tr>
<td>History of ICH</td>
<td>2 (20.0)</td>
<td>1 (2.6)</td>
<td>0.10</td>
</tr>
<tr>
<td>ICH volume, mL ± SD</td>
<td>31.2±35.6</td>
<td>43.9±24.0</td>
<td>0.19</td>
</tr>
<tr>
<td>nWMH</td>
<td>17.3 [9.3, 31.6]</td>
<td>15.9 [2.8, 33.8]</td>
<td>0.67</td>
</tr>
<tr>
<td>No. of MB</td>
<td>1 [0, 4]</td>
<td>0.5 [0, 3]</td>
<td>0.76</td>
</tr>
<tr>
<td>Mean ADC, $10^{-4}$ mm$^2$/s</td>
<td>12.4±0.89</td>
<td>11.7±0.91</td>
<td>0.03</td>
</tr>
<tr>
<td>Visible cerebral atrophy score</td>
<td>5.1±3.6</td>
<td>3.0±3.1</td>
<td>0.08</td>
</tr>
</tbody>
</table>

Values are displayed as mean ± SD, median [25th and 75th quartile], or n (%).
tion before CAA-related ICH is reported to be 20% to 40%.5,17 Autopsy-based studies of stroke-free individuals show that CAA is a risk factor for decreased antemortem cognitive performance while simultaneously controlling for the pathology of Alzheimer disease.31,32 This suggests that the association between CAA and cognitive dysfunction is not entirely mediated by concomitant Alzheimer disease. Another plausible mechanism by which CAA might cause cognitive dysfunction is impaired vascular dysfunction with intermittent ischemia.3 White matter damage, a marker of vascular disease,33–35 is seen in CAA4,14 and has been associated with PICI.4 The lack of association between nWMH volume and PICI in the current study likely reflects the small sample size and highlights the robustness of the association detected between mean ADC and PICI.

Global mean ADC measures the degree of chronic tissue disruption and has been shown, in other cerebral small vessel diseases, to be a very sensitive measure of cerebral tissue damage.9,11,13 The strong independent association between mean ADC and PICI that we observed in this study suggests that global MRI diffusion changes may be a useful marker of CAA-related cognitive impairment by detecting clinically relevant tissue microstructural alterations. These microstructural alterations could be caused by direct ischemic injury or decreased white matter organization from neurodegeneration with neuronal loss.9,11,13

In a previous study, we found an association between nWMH volume and subcortical regional fractional anisotropy measures,12 whereas in this work, we did not find an association between nWMH volume and mean ADC. This may be related to our use of global measures, instead of region of interest-based measures of fractional anisotropy from subcortical regions, or may be related to the use of ADC rather than fractional anisotropy. Interestingly, in our previous study comparing subjects with CAA and control subjects, fractional anisotropy differences were greatest in regions of white matter and gray matter not typically involved by WMH.12 These observations are consistent with previous studies in sporadic leukoaraiosis16,37 and cerebral autosomal-dominant arteriopathy with subcortical ischemic leukoencephalopathy9 that show that diffusion tensor imaging measures are sensitive to tissue changes in apparently nonlesional areas on T2 sequences.

The main limitation of this study is the relatively small sample size, which may have precluded the detection of independent associations of MRI markers, other than global mean ADC, with PICI. The fact that global mean ADC was the only MRI marker associated with PICI attests to the strength of the observed association, however. Other vascular risk factors may act synergistically to cause tissue microstructural changes in patients with CAA, although we were unable to detect any significant associations between these variables and mean ADC in this small cohort. Further studies are necessary to confirm the findings seen in the current work. Second, because data from clinical MRI images were analyzed in this study, the spatial resolution was insufficient to perform quantitative volumetric analyses of brain atrophy. Our semiquantitative measure may not fully account for the impact on cerebral atrophy on PICI. Further studies using serial high-resolution quantitative MRI are ongoing. We cannot definitively exclude that some subjects had ICH from non-CAA-related pathology; however, such subjects likely represent a small portion of the cohort. The likelihood of underlying CAA was assessed using the validated, highly specific Boston criteria.16,24 The majority of subjects with PICI (7 of 10) had a diagnosis of probable CAA according to these criteria; this diagnosis has near 100% specificity for the pathological presence of CAA.16

In summary, these results show that the degree of tissue microstructural alteration, quantified by global mean ADC, is strongly associated with pre-ICH cognitive impairment independent of age and other MRI markers, including a measure of brain atrophy. The lack of correlation with most other MRI markers suggests that mean ADC may be sensitive to distinct aspects of CAA pathology and its tissue consequences. Further prospective studies in CAA will attempt to define the relationship among changes in mean ADC, other MRI markers, and clinical impairment. More generally, these results as well as other studies in cerebral autosomal-dominant arteriopathy with subcortical ischemic leukoencephalopathy11,13 suggest that mean ADC should be evaluated as a potential marker of cognitive dysfunction in MRI studies of cerebral small vessel diseases and cognition.

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**Disclosures**

None.

**References**


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