Hippocampal Volumes, Proton Magnetic Resonance Spectroscopy Metabolites, and Cerebrovascular Disease in Mild Cognitive Impairment Subtypes

Kejal Kantarci, MD; Ronald C. Petersen, MD, PhD; Scott A. Przybelski, BS; Stephen D. Weigand, MS; Maria M. Shiung, BA; Jennifer L. Whitwell, PhD; Selamawit Negash, PhD; Robert J. Ivnik, PhD; Bradley F. Boeve, MD; David S. Knopman, MD; Glenn E. Smith, PhD; Clifford R. Jack Jr, MD

Background: Although a majority of patients with amnestic mild cognitive impairment (aMCI) progress to Alzheimer disease, the natural history of nonamnestic MCI (naMCI) is less clear. Noninvasive imaging surrogates for underlying pathological findings in MCI would be clinically useful for identifying patients who may benefit from disease-specific treatments at the prodromal stage of dementia.

Objective: To determine the characteristic magnetic resonance imaging (MRI) and proton MR spectroscopy (1H MRS) profiles of MCI subtypes.

Design: Case-control study.

Setting: Community-based sample at a tertiary referral center.

Patients: Ninety-one patients with single-domain aMCI, 32 patients with multiple-domain aMCI, 20 patients with single- or multiple-domain naMCI, and 100 cognitively normal elderly subjects frequency-matched by age and sex.

Main Outcome Measures: Posterior cingulate gyrus 1H MRS metabolite ratios, hippocampal volumes, and cerebrovascular disease on MRI.

Results: Patients with single-domain aMCI were characterized by small hippocampal volumes and elevated ratios of myo-inositol to creatine levels. Patients with naMCI on average had normal hippocampal volumes and 1H MRS metabolite ratios, but a greater proportion (3 of 20 patients [15%]) had cortical infarctions compared with patients with single-domain aMCI (6 of 91 [7%]). For characterization of MCI subtypes, 1H MRS and structural MRI findings were complementary.

Conclusions: The MRI and 1H MRS findings in single-domain aMCI are consistent with a pattern similar to that of Alzheimer disease. Absence of this pattern on average in patients with naMCI suggests that cerebrovascular disease and other neurodegenerative diseases may be contributing to the cognitive impairment in many individuals with naMCI.

Arch Neurol. 2008;65(12):1621-1628

THE BROAD CLINICAL DEFINITION of mild cognitive impairment (MCI) includes amnestic MCI (aMCI) and nonamnestic MCI (naMCI), with impairment in a single cognitive domain or multiple cognitive domains such as memory, attention/executive functioning, language, and visuospatial processing.1 Although patients with the aMCI subtype have a higher risk of progression to Alzheimer disease (AD) compared with their cognitively normal peers, the natural history and the pathological underpinnings of naMCI are less clear.2-8 However, longitudinal studies suggest that patients with naMCI also have a higher rate of progression to dementia than do cognitively normal individuals.2-5,8 The most common pathological findings encountered in aMCI include AD, cerebrovascular disease, and Lewy body pathological features.9-13 Noninvasive imaging surrogates for these underlying abnormalities in MCI would be clinically useful for identifying patients who may benefit from disease-specific treatments at the prodromal stage of dementia.

Hippocampal volumetry based on magnetic resonance imaging (MRI) and proton magnetic resonance spectroscopy (1H MRS) metabolite ratios are sensitive markers for AD pathological changes.14-20 Cortical and subcortical infarctions represent cerebrovascular disease, and evidence exists that white matter hyperintensities (WMHs) are related to ischemic vascular disease.18 In cross-sectional 1H MRS studies, the glial activation marker myo-inositol (ml) or the ratio of ml to creatine (ml:Cr ratio) and the membrane integrity marker choline (Cho) or the Cho:Cr ratio are elevated in aMCI, AD, and dementia.
with Lewy bodies (DLB). The level of the neuronal integrity marker N-acetylaspartate (NAA) tends to be decreased in patients with aMCI, AD, and vascular dementia.21-28 Whereas aMCI has been the focus of much research in identifying imaging markers for prodromal AD, few studies have investigated the imaging characteristics of naMCI.29-33

Because of the complexity of pathological findings underlying dementia, it is conceivable that magnetic resonance markers that are surrogates for various pathological substrates of dementia may provide complementary information concerning the typical underlying causes of dementia in the elderly. Our objective was to determine the characteristic magnetic resonance profiles of the MCI subtypes, including hippocampal volumes, \(^1\)H MRS metabolite ratios, and cerebrovascular disease.

### METHODS

**RECRUITMENT OF SUBJECTS**

We identified 91 patients with single-domain aMCI, 32 patients with multiple-domain aMCI, and 20 patients with single- or multiple-domain naMCI who were consecutively recruited to the Mayo Clinic AD Research Center (ADRC) and AD patient registry (ADPR) cohorts and who participated in an MRI-\(^1\)H MRS study. From the same ADRC-ADPR cohort, we identified 100 cognitively normal subjects who were frequency-matched by age and sex to the MCI patients and who underwent MRI during the same period. This study was approved by the Mayo Clinic institutional review board, and informed consent for participation was obtained from every subject and/or an appropriate surrogate.

Individuals participating in the ADRC-ADPR studies underwent approximately annual clinical examinations, structural brain MRI and \(^1\)H MRS, routine laboratory tests, and a battery of neuropsychological tests. At the completion of the evaluation, a consensus committee meeting is held involving the behavioral neurologists (R.C.P., B.F.B., and D.S.K.), neuropsychologists (R.J.I. and G.E.S.), nurses, and the geriatrician who performed the evaluations of the subjects to assign a clinical diagnosis to the participant.

The operational definition of MCI was based on clinical judgment through a careful history obtained from the patient and preferably a collateral source without reference to MRI results, using the following criteria for the broad definition of MCI: cognitive complaint, cognitive function not normal for age, decline in cognition, essentially normal functional activities, and no dementia. The patients with MCI were further classified into 1 of the following 4 MCI subtypes: (1) single-domain aMCI, if the impairment was only in the memory domain; (2) multiple-domain aMCI, if the impairment was in the memory domain plus 1 or more other domains such as language, attention/executive function, and visuospatial processing; (3) single-domain naMCI, if the impairment was in 1 or more nonmemory domains with relative preservation of memory; and (4) multiple-domain naMCI, if the impairment was in more than 1 domain with relative preservation of memory. We grouped patients with naMCI and impairment in single and multiple domains in this study because of the small number of patients with multiple-domain naMCI. We excluded patients with structural abnormalities that could impair cognitive function other than cerebrovascular lesions, such as tumor, subdural hematoma, and contusion due to a previous head trauma, and patients with addictions, psychiatric diseases, or treatments that would affect cognitive function. Subjects were not excluded for the presence of infarctions and leukoaraiosis; thus, the full range of ischemic cerebrovascular disease was included.

Cognitively normal subjects were recruited from the community but underwent evaluation in the same manner as patients with MCI. The cognitively normal group did not have any neurological or psychiatric conditions, did not have a cognitive concern, had normal results of the neurological and neuropsychological examinations, and were not taking psychoactive medications in doses that would affect cognition. To ensure that the cognitively normal group did not include subjects with preclinical dementia, we included subjects only if they remained cognitively normal for at least 2 years during longitudinal clinical follow-up.

Apolipoprotein E genotype is determined by established polymerase chain reaction techniques. Participants with genotypes 2/4, 3/4, and 4/4 were labeled ε4 carriers; those with the other genotypes were labeled ε4 noncarriers.

### NEUROPSYCHOLOGICAL TESTING

Cognitive testing was completed within 4 months of the MRI and \(^1\)H MRS studies. Memory was evaluated by means of freerecall percentage of retention scores computed after a 30-minute delay for the Wechsler Memory Scale–Revised logical memory and visual reproduction subtests and the Rey Auditory Verbal Learning Test.34 Language tests measured naming to confrontation (ie, the Boston Naming Test)35 and category fluency (ie, naming animals, fruits, and vegetables).36 The attention/executive measures included the Trail-Making Test parts A and B36 and the Wechsler Adult Intelligence Scale–Revised digit symbol subtest. Visuospatial processing was examined by means of the picture completion and block design subtests of the Wechsler Adult Intelligence Scale–Revised. All tests were administered by experienced psychometrists and supervised by clinical neuropsychologists (R.J.I. and G.E.S.). All raw scores were converted to Mayo Older Americans Normative Studies (MOANS) age-adjusted scaled scores that are normally distributed and that have a mean (SD) score of 10 (3) in cognitively healthy subjects, on whom each test was normed.37,39,40 In each cognitive domain, we computed a mean MOANS age-corrected scaled score for every participant. We note that the patients’ mean MOANS scores within a certain domain did not strictly define their MCI category, although cognitive tests were used to inform the clinical consensus diagnosis.

### MRI AND \(^1\)H MRS STUDIES

All subjects underwent MRI and \(^1\)H MRS studies on a 1.5-T scanner (Signa; GE Medical Systems, Milwaukee, Wisconsin). Hippocampal volume measurements were derived from a T1-weighted 3-dimensional volumetric spoiled gradient-echo sequence with 124 continuous partitions, 1.6-mm section thickness, a 22 × 16.5-cm or 24 × 18.5-cm field of view, 192 views, and 25° flip angle. Hippocampal volumes were measured by manually tracing their anatomical boundaries for each image section sequentially from posterior to anterior.41 The intrarater test-retest coefficient of variation of hippocampal volume measurement is 0.97% for the image analyst (M.M.S.), who manually traced each hippocampus and was masked to all clinical information. Volumes were then converted to normal deviates, referred to as W scores, using age- and sex-specific normal percentiles based on a previous study.42 Among cognitively normal subjects, a value of 0 corresponds to the 50th percentile; 1.64, to the 95th percentile; and –1.64, to the 5th percentile.

A fluid-attenuated inversion recovery (FLAIR) pulse sequence (repetition time, 16 000 milliseconds; time following...
inversion pulse, 2600 milliseconds; echo time, 140 milliseconds; 256 × 160 matrix; 1 repetition; 22-cm field of view; and 3-mm interleaved images of the whole head) was used for the assessment of cerebrovascular disease. A radiologist (K.K.) blinded to all clinical images assessed the WMHs and hemispheric cortical and lacunar infarctions. The WMH volume was estimated by visually comparing the subject’s FLAIR images with a bank of 10 FLAIR image templates with increasing WMH volumes (from 1 to 100 cm³) determined with an automated image segmentation algorithm.44 A continuous scale with a slider bar was used to estimate each subject’s WMH volume by matching to the WMH templates.44 The WMH volume estimation algorithm was previously validated against quantification using the automated image segmentation of the WMH volume (concordance correlation coefficient, 0.88; 95% confidence interval, 0.83-0.94). Intrarater reliability for assessment of the WMHs is excellent (intraclass correlation coefficient, 0.98; 95% confidence interval, 0.97-0.98). Hemispheric cortical infarctions were defined as areas of elevated signal intensity involving the cortical gray mantle and immediately subjacent white matter that exceed 1 cm in the largest diameter on FLAIR images. Subcortical infarctions were defined as discrete subcortical lesions of more than 3 mm in diameter with intensity that is equivalent to cerebrospinal fluid on FLAIR images and accompanying hyperintense gliotic rim. Intrarater reliability for assessment of subcortical infarctions (proportion in agreement, 0.94) and cortical infarctions (proportion in agreement, 0.98) is excellent.

The T1-weighted images in the sagittal plane were obtained for localizing the 1H MRS voxel. The 1H MRS studies were performed with an automated single-voxel MRS package (Proton Brain Examination/Single Voxel; GE Medical Systems).60 Point-resolved spectroscopy pulse sequences (repetition time, 2000 milliseconds; echo time, 30 milliseconds; 2048 data points; and 128 excitation) were used for the examinations. An 8-cm³ (2 × 2 × 2-cm) voxel, prescribed on a midsagittal T1-weighted image, included the right and left posterior cingulate gyri and inferior precunei.60 We analyzed the metabolite intensity ratios, using Cr as an internal reference metabolite.

STATISTICAL ANALYSIS

We compared the cognitively normal subjects and patients with the 3 MCI subtypes on categorical patient characteristics and MRI findings such as the presence of cortical and subcortical infarctions using the χ² test, or, if expected cell counts were less than 5, the Fisher exact test. We used Kruskal-Wallis tests to compare groups on age, education, MOANS scores for the 4 cognitive domains, and quantitative magnetic resonance measurements. Pairwise group comparisons among the clinical groups on quantitative imaging measures were performed using Wilcoxon rank sum tests because of skewness. These tests were all 2 sided.

To assess the effect of imaging measures on the estimated probability of cognitively normal, single-domain aMCI, multiple-domain aMCI, or naMCI findings, we used generalized logit multinominal models.47 These models extend binary logistic regression to the situation where the response is an unordered categorical variable with 3 or more levels. Models included age, sex, and education as adjustment variables and examined the effects of a single imaging predictor. We performed a likelihood ratio test to evaluate whether adding the imaging predictor significantly improved model fit (and therefore discrimination) compared with a model with age, sex, and education only (ie, a model without imaging information). We graphically illustrated the effect of the imaging predictor on the estimated probability of group membership by showing relative probabilities as a function of the imaging predictor. The relative probability is the estimated probability of group membership at a given imaging value divided by the estimated probability at the median or at these estimates. We arbitrarily set sex to male, age to 77 years, and education to 14 years, with covariate levels representing the most common category or the median value in our sample.

We also used multinomial modeling to fit what we call 2-imaging predictor models that included age, sex, education, and hippocampal W score plus 1 of the following: NAA:Cr ratio, ml:Cr ratio, Cho:Cr ratio, the log of WMH, and cortical infarctions. These models were used to evaluate whether a measurement improved model fit and discrimination beyond that obtained from an age-, sex-, and education-adjusted model based on hippocampal volume only. The significance of the second imaging predictor was evaluated using a likelihood ratio test. We did not adjust for multiple comparisons because the statistical tests address questions of distinct albeit related clinical interest.59 60 We used commercially available software (SAS, version 8.2 [SAS Institute Inc, Cary, North Carolina] and R, version 2.7.0 [R Foundation for Statistical Computing, Vienna, Austria]) for these analyses.

RESULTS

Cognitively normal subjects and patients with MCI did not differ significantly in age, sex, and years of education. As expected, a greater proportion of patients with MCI than cognitively normal subjects were apolipoprotein E ε4 carriers. Table 1 demonstrates that, although general cognitive function as indicated by Mini-Mental State Examination and Clinical Dementia Rating sum of boxes scores are identical in terms of median value across the MCI subtypes, the specific domains that are impaired as indicated by lower MOANS scores differ significantly by MCI subtype.

The patients with single-domain aMCI had smaller hippocampal W scores than did cognitively normal subjects (P < .001) and patients with multiple-domain aMCI (P = .009) and naMCI (P = .008). Similarly, patients with single-domain aMCI had elevated ml:Cr ratios on 1H MRS compared with cognitively normal subjects (P < .001) and patients with multiple-domain aMCI (P = .02) and naMCI (P = .03). Patients with multiple-domain aMCI had a trend toward smaller hippocampal W scores (P = .046) and lower NAA:Cr ratios on 1H MRS (P = .06) compared with cognitively normal subjects. Patients with naMCI had normal hippocampal volumes and 1H MRS metabolite ratios (Table 2).

A greater observed proportion of patients with multiple-domain aMCI (16%) and with naMCI (15%) had cortical infarctions compared with patients with single-domain aMCI (7%). Although the proportion of the patients with cortical infarctions was twice as high in multiple-domain aMCI and naMCI compared with single-domain aMCI, this difference was statistically a trend (P = .17). If the true rates of cortical infarctions were 0.07 in the single-domain aMCI group, 0.16 in the multiple-domain aMCI group, and 0.15 in the naMCI group (ie, what we observed), then we would have approximately 38% power to detect a significant difference using the Fisher exact test. On the other hand, the proportion of patients with subcortical infarctions was similar across
Table 1. Demographic Characteristics and Cognitive Features of the Cohort

<table>
<thead>
<tr>
<th>Subject Groups</th>
<th>CN (n=100)</th>
<th>SD-aMCI (n=91)</th>
<th>MD-aMCI (n=32)</th>
<th>naMCI (n=20)</th>
<th>P Valueb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Women, No. (%)</td>
<td>42 (42)</td>
<td>40 (44)</td>
<td>14 (44)</td>
<td>7 (35)</td>
<td>.90</td>
</tr>
<tr>
<td>Age, median (IQR), y</td>
<td>76 (71-81)</td>
<td>78 (71-93)</td>
<td>78 (72-83)</td>
<td>76 (66-80)</td>
<td>.17</td>
</tr>
<tr>
<td>Education, y</td>
<td>14 (12-16)</td>
<td>15 (12-18)</td>
<td>13 (12-16)</td>
<td>12 (12-15)</td>
<td>.28</td>
</tr>
<tr>
<td>APOE ε4 carriers, No. (%)</td>
<td>26 (26)</td>
<td>41 (45)</td>
<td>13 (41)</td>
<td>8 (40)</td>
<td>.01</td>
</tr>
<tr>
<td>Memory MOANS score</td>
<td>11.0 (10.3-12.0)</td>
<td>6.4 (4.8-7.8)</td>
<td>7.7 (5.9-8.5)</td>
<td>9.1 (8.3-10.1)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Language MOANS score</td>
<td>10.8 (9.7-12.7)</td>
<td>9.3 (8.0-11.0)</td>
<td>7.8 (6.3-9.3)</td>
<td>8.8 (7.2-10.0)</td>
<td>.002</td>
</tr>
<tr>
<td>Attention/executive MOANS score</td>
<td>11.0 (10.3-12.0)</td>
<td>9.8 (8.7-11.0)</td>
<td>7.3 (6.3-9.0)</td>
<td>7.8 (6.7-9.4)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Visuospatial processing MOANS score</td>
<td>11.0 (9.8-12.5)</td>
<td>10.5 (9.0-12.0)</td>
<td>7.0 (6.5-9.0)</td>
<td>8.5 (6.8-10.0)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>MMSE</td>
<td>29 (28-30)</td>
<td>27 (25-29)</td>
<td>27 (25-28)</td>
<td>27 (26-29)</td>
<td>.42</td>
</tr>
<tr>
<td>GDR sum of boxes</td>
<td>0 (0-0)</td>
<td>0.5 (0.5-1.5)</td>
<td>0.5 (0.5-1.0)</td>
<td>0.5 (0.5-2.0)</td>
<td>.92</td>
</tr>
</tbody>
</table>

Abbreviations: APOE, apolipoprotein E; CDR, Clinical Dementia Rating; CN, cognitively normal; IQR, interquartile range; MD-aMCI, multiple-domain amnestic mild cognitive impairment (MCI); MOANS, Mayo Older Americans Normative Studies; naMCI, nonamnestic MCI; SD-aMCI, single-domain aMCI.

a Except where indicated, values shown are median (interquartile range).
b P values for sex, age, education, and APOE genotype compare all 4 groups, whereas P values for cognitive variables compare MCI subtypes. Sex and APOE genotype are tested using a χ² test, whereas other variables are tested with a Kruskal-Wallis test.

Table 2. MRI and 1H MRS Findings in Cognitively Normal Subjects and MCI Subtypes

<table>
<thead>
<tr>
<th>Subject Groups</th>
<th>CN (n=100)</th>
<th>SD-aMCI (n=91)</th>
<th>MD-aMCI (n=32)</th>
<th>naMCI (n=20)</th>
<th>P Valueb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hippocampal W score</td>
<td>0.29 (-0.50 to 0.74)</td>
<td>-1.01 (-1.93 to -0.01)</td>
<td>-0.38 (-0.73 to 0.58)</td>
<td>0.21 (-0.85 to 0.75)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>NAA:Cr ratio</td>
<td>1.52 (1.46 to 1.59)</td>
<td>1.51 (1.43 to 1.58)</td>
<td>1.50 (1.42 to 1.53)</td>
<td>1.53 (1.44 to 1.58)</td>
<td>.28</td>
</tr>
<tr>
<td>ml:Cr ratio</td>
<td>0.67 (0.62 to 0.71)</td>
<td>0.70 (0.66 to 0.74)</td>
<td>0.67 (0.61 to 0.72)</td>
<td>0.65 (0.62 to 0.72)</td>
<td>.003</td>
</tr>
<tr>
<td>Cho:Cr ratio</td>
<td>0.64 (0.60 to 0.70)</td>
<td>0.66 (0.64 to 0.73)</td>
<td>0.68 (0.63 to 0.72)</td>
<td>0.67 (0.60 to 0.74)</td>
<td>.012</td>
</tr>
<tr>
<td>WMH volume, cm³</td>
<td>10 (6 to 17)</td>
<td>16 (6 to 23)</td>
<td>10 (7 to 23)</td>
<td>14 (7 to 30)</td>
<td>.02</td>
</tr>
<tr>
<td>No. (%) of subjects with any subcortical infarctions</td>
<td>10 (10)</td>
<td>9 (10)</td>
<td>3 (9)</td>
<td>2 (10)</td>
<td>&gt;.99</td>
</tr>
<tr>
<td>No. (%) of subjects with any cortical infarctions</td>
<td>0</td>
<td>6 (7)</td>
<td>5 (16)</td>
<td>3 (15)</td>
<td>.17</td>
</tr>
</tbody>
</table>

Abbreviations: CN, cognitively normal; Cho, choline; Cr, creatine; 'H MRS, proton magnetic resonance spectroscopy; MCI, mild cognitive impairment; MD-aMCI, multiple-domain amnestic MCI; ml, myo-inositol; MRI, magnetic resonance imaging; NAA, N-acetylaspartate; naMCI, nonamnestic MCI; SD-aMCI, single-domain aMCI; WMH, white matter hyperintensity.

a Except where indicated, values shown are median (interquartile range).
b P values for continuous imaging measures are tested using a Kruskal-Wallis test among all 4 groups, whereas infarction variables are tested using a Fisher exact test among MCI subtypes only.

MCI subtypes and cognitively normal subjects. The average estimated volume of WMH in patients with single-domain aMCI was higher than in cognitively normal subjects (P = .002). Despite the fact that, on average, the patients with multiple-domain aMCI had lower WMH volumes than did those with single-domain aMCI and naMCI subtypes, the 3 MCI groups did not differ on WMH volume, which may be a result of the small sample sizes of the multiple-domain aMCI and naMCI groups (Table 2).

The Figure illustrates that among the cognitively normal subjects and MCI subtypes, after adjusting for age, sex, and education, the estimated probability of being in the single-domain aMCI group was higher at lower hippocampal W scores and higher ml:Cr ratios than in the cognitively normal subjects and the patients with naMCI (generalized logistic multinomial models, P < .001). Whereas the estimated probabilities in single-domain aMCI and naMCI were in general in the opposite direction of those for the cognitively normal individuals, quantitative imaging markers did not seem to affect the probability of membership in the multiple-domain aMCI group.

Multinomial modeling to fit the 2 imaging-predictor models that included age, sex, education, and a single imaging marker were used to evaluate whether a second imaging measurement improved model fit and discrimination beyond that obtained from an age-, sex-, and education-adjusted model based on a single imaging marker only. We included cognitively normal subjects and each of the MCI subtypes in the first model and included only the MCI subtypes in the second model to determine whether MRI and 'H MRS variables were complementary in distinguishing MCI subtypes. Higher ml:Cr and Cho:Cr ratios and greater WMH volume improved the discrimination among all clinical groups (MCI subtypes and cognitively normal subjects) when controlled for hippocampal W scores or other imaging markers, demonstrating that hippocampal W scores, ml:Cr and Cho:Cr ratios, and WMHs are complementary in characterizing patients with MCI and cognitively normal subjects. On the other hand, only higher ml:Cr ratios improved the discrimination among the MCI subtypes when we controlled for hippocampal W scores and WMH volume (Table 3).
There were distinct groupwise differences in MRI and $^1$H MRS findings between single-domain aMCI and naMCI subtypes. Patients with single-domain aMCI tended to have smaller hippocampal volumes and elevated mI:Cr ratios compared with patients with naMCI and cognitively normal subjects. On the other hand, patients with naMCI had normal hippocampal volumes and normal mI:Cr ratios, but a greater proportion of these patients in our sample had cortical infarctions compared with the patients with single-domain aMCI. Both hippocampal atrophy and elevated mI:Cr ratios are sensitive markers of early AD pathological changes, and the severity of these abnormalities correlate with pathological severity of AD. For this reason, hippocampal atrophy and elevated mI:Cr ratios most likely represent a high frequency of early AD pathological changes in the patients with single-domain aMCI. On average, hippocampal volumes and mI:Cr ratios in the naMCI subtype suggest that underlying pathological substrates may include abnormalities other than AD in some patients with naMCI. Mixed brain abnormalities, including AD cerebrovascular and Lewy body disease, underlie most cases of dementia in the community. It appears likely from the higher prevalence of cortical infarctions in patients with naMCI that large-vessel cerebrovascular disease was one of the pathological contributors to naMCI, in agreement with a previous study showing that a greater proportion of patients with naMCI experienced transient ischemic attacks and stroke compared with patients with aMCI. On the other hand, we found a similar apolipoprotein E ε4 frequency in the naMCI and aMCI groups, yet we dem-

Figure. Estimated relative probability of group membership as a function of 4 imaging measures. The relative probabilities shown are the estimated probability of group membership obtained from the multinomial model divided by the estimated probability at the median of the imaging measure. The gray reference lines intersect at the median for the measure on the x-axis and 1.0 on the y-axis. For all estimates we assumed a 77-year-old man with 14 years of education. Cho indicates choline; CN, cognitively normal; Cr, creatine; MD-aMCI, multiple-domain amnestic mild cognitive impairment (MCI); mI, myo-inositol; naMCI, nonamnestic MCI; SD-aMCI, single-domain aMCI; and WMH, white matter hyperintensity.
onstrate that patients with naMCI on average do not have the magnetic resonance features of AD, and by definition they present with an early AD phenotype (ie, predominantly memory impairment). One possible way to reconcile these seemingly contradictory findings, although speculative, is that some patients with naMCI have AD pathological features but in an atypical anatomical distribution, thus accounting for the non–AD-like magnetic resonance findings and clinical presentation. In support of this, autopsy studies have documented the presence of AD pathological characteristics in an atypical distribution in subjects who present with a non–AD-like clinical profile.\(^3^1\)\(^3^2\) However, this will require follow-up, preferably to autopsy.

Multinominal modeling demonstrated that elevated ml:Cr and Cho:Cr ratios and increased WMH volumes were complementary to smaller hippocampal volumes in distinguishing the clinical groups. This is in agreement with a previous finding that \(^1\)H MRS metabolites improve discrimination of cognitively impaired nondemented individuals from their cognitively normal peers, when considered together with hippocampal volumes.\(^2^3\) Furthermore, \(^1\)H MRS findings and hippocampal volume are independent and complementary predictors of verbal memory on neuropsychometric testing in nondemented older adults.\(^5^3\) The added value of \(^1\)H MRS metabolites, hippocampal volumes, and WMH volumes in characterizing the MCI subtypes suggests that \(^1\)H MRS scans may complement structural MRI findings as a prognostic marker in MCI; however, this will require longitudinal studies for verification.

The magnetic resonance findings in patients with multiple-domain aMCI showed some similarities to those of patients with single-domain aMCI and naMCI. These patients had some hippocampal atrophy, but this was less than the hippocampal atrophy observed in patients with single-domain aMCI. Furthermore, they typically had normal ml:Cr ratios on \(^1\)H MRS findings, suggesting that if these patients had pathological AD, it would be on average less severe than in patients with single-domain aMCI. On the other hand, in our sample, the proportion of patients with multiple-domain aMCI and cortical infarctions was more than twice as high as that for patients with single-domain aMCI. For this reason, our data suggest that AD and cerebrovascular disease, in variable amounts, are some of the underlying abnormalities in the patients with multiple-domain aMCI in this study.

<table>
<thead>
<tr>
<th>MCI Subtype</th>
<th>Hippocampal Volumes</th>
<th>(^1)H MRS, ml:Cr Ratio</th>
<th>Cortical Infarctions</th>
<th>Pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single-domain aMCI</td>
<td>Atrophy</td>
<td>Elevated</td>
<td>Less frequent</td>
<td>Yes</td>
</tr>
<tr>
<td>naMCI</td>
<td>Normal</td>
<td>Normal</td>
<td>More frequent</td>
<td>No</td>
</tr>
<tr>
<td>Multiple-domain aMCI</td>
<td>Mild atrophy</td>
<td>Normal</td>
<td>More frequent</td>
<td>Some</td>
</tr>
</tbody>
</table>

Abbreviations: AD, Alzheimer disease; aMCI, amnestic mild cognitive impairment (MCI); Cr, creatine; CVD, cerebrovascular disease; \(^1\)H MRS, proton magnetic resonance spectroscopy; ml, myo-inositol; MRI, magnetic resonance imaging; naMCI, nonamnestic MCI.
Contrary to the evidence of varying prevalence rates of cortical infarctions, we did not observe any difference in the proportion of patients with subcortical infarctions among cognitively normal subjects and the 3 MCI subtypes. The possibility that cortical and subcortical infarctions have a differential effect on future progression to dementia is currently unclear. One study found that subcortical infarctions did not increase the risk of progression to dementia in MCI. Others have shown that silent infarctions, most of which were presumably subcortical, increase the risk of future dementia in cognitively normal elderly individuals. The patients with single-domain aMCI had a greater WMH volume compared with cognitively normal elderly subjects. On the other hand, we did not identify a difference in the estimated WMH volume among the MCI subtypes. Because the frequency of subcortical infarctions was also similar across the MCI subtypes, our data suggest that subcortical vascular disease is a common but not a differentiating feature of the MCI subtypes. This is consistent with the previous reports that WMH is associated with an increased risk of MCI and that there is no association between subcortical hyperintensity load and different MCI subtypes.

This study has several limitations. First, we classified infarctions as cortical and subcortical and did not further investigate the effects of the location, number, and size of the infarctions on cognitive function in MCI. This was a deliberate choice because the small number of subjects with infarctions prevented any coherent grouping of infarctions by detailed anatomical criteria for groupwise analysis. Second, cognitively normal subjects were neurologically healthy individuals who did not have a clinical history of stroke or cortical infarctions. They were included in this study to test for the abnormalities in quantitative magnetic resonance markers in MCI subtypes without the confounding effects of overt known neurological diseases. It is possible, although not common, for cognitively normal subjects to have cortical infarctions. Therefore, the absence of cortical infarctions in the cognitively normal subjects of this study cannot be generalized to the population. Third, although it is expected, based on pathological series, that DLB is one of the underlying abnormalities in MCI subtypes, the presence of DLB-related imaging changes was not evaluated in this study because there are no established magnetic resonance markers for prodromal DLB. Normal hippocampal volumes and $^1$H MRS in naMCI do not rule out the presence of Lewy body disease because patients with DLB on average have normal hippocampal volumes and normal posterior cingulate gyrus NAA:Cr and mI:Cr ratios. Although patients with DLB have elevated Cho:Cr ratios on $^1$H MRS findings, it is not clear whether the Cho:Cr ratio elevation in the posterior cingulate gyrus is a feature of prodromal DLB. For these reasons, DLB may be one of the underlying pathological findings in MCI subtypes.

Our data indicate that specific constellations of magnetic resonance findings are complementary in characterizing MCI subtypes and cognitively normal individuals. The MRI and $^1$H MRS findings in patients with single-domain aMCI are characterized by an AD-like pattern of elevated mI:Cr ratios in the posterior cingulate gyrus and hippocampal atrophy. On the other hand, patients with naMCI do not have the magnetic resonance features of AD and are more likely to have cortical infarctions. Magnetic resonance findings in patients with multiple-domain aMCI show similarities to those in patients with single-domain aMCI and naMCI. Clinical follow-up will demonstrate the combinations of MRI and $^1$H MRS findings that are useful in determining which MCI patients will experience progression to specific dementia syndromes in the future.

Accepted for Publication: August 7, 2008.
Correspondence: Kejal Kantarci, MD, Department of Radiology, Mayo Clinic, 200 First St SW, Rochester, MN 55905 (kantarci.kejal@mayo.edu).
Author Contributions: Dr Kantarci had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. Study concept and design: Kantarci, Whitwell, and Jack. Acquisition of data: Kantarci, Shiung, Whitwell, Ivnik, Negash, Boeve, and Jack. Analysis and interpretation of data: Kantarci, Petersen, Przybelski, Weigand, Negash, Boeve, and Jack. Drafting of the manuscript: Kantarci. Critical revision of the manuscript for important intellectual content: Petersen, Przybelski, Weigand, Shiung, Whitwell, Negash, Ivnik, Boeve, Knopman, Smith, and Jack. Statistical analysis: Kantarci, Przybelski, and Weigand. Obtained funding: Kantarci, Petersen, and Jack. Administrative, technical, and material support: Kantarci, Shiung, Ivnik, Boeve, Smith, and Jack. Study supervision: Jack.
Financial Disclosure: None reported.
Funding/Support: This study was supported by Paul B. Beeson Career Development Award in Aging K23 AG030935, from the National Institutes of Health (NIH)/National Institute on Aging (NIA); Alzheimer’s Association New Investigator Research Grant 03-4842; NIH Roadmap Multidisciplinary Clinical Research Career Development awards KL2 RR024151 from the NIH/National Center for Research Resources (Dr Kantarci), P50 AG16574 and U01 AG06786 from the NIH/NIA (Dr Petersen), and RO1 AG11378 from the NIH/NIA (Dr Jack); the Alexander Family Professorship in Alzheimer’s Research (Dr Jack); and the Robert H. and Clarice Smith and Abigail Van Buren Alzheimer’s Disease Research Program.

REFERENCES
