In Vivo Evaluation of Brain Iron in Alzheimer Disease Using Magnetic Resonance Imaging

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Background: The basal ganglia contain the highest levels of iron in the brain, and postmortem studies indicate a disruption of iron metabolism in the basal ganglia of patients with Alzheimer disease (AD). Iron can catalyze free radical reactions and may contribute to oxidative damage observed in AD brains. Treatments aimed at reducing oxidative damage have offered novel ways to delay the rate of progression and could possibly defer the onset of AD. Brain iron levels were quantified in vivo using a new magnetic resonance imaging method.

Methods: Thirty-one patients with AD and 68 control subjects participated in this study. A magnetic resonance imaging method was employed that quantifies the iron content of ferritin molecules (ferritin iron) with specificity through the combined use of high and low field-strength magnetic resonance imaging instruments. Three basal ganglia structures (caudate, putamen, and globus pallidus) and one comparison region (frontal lobe white matter) were evaluated.

Results: Basal ganglia ferritin iron levels were significantly increased in the caudate ($P = .007$; effect size, 0.69) and putamen ($P = .008$; effect size, 0.67) of AD subjects, with a trend toward an increase in the globus pallidus ($P = .13$). The increased basal ganglia ferritin iron levels were not a generalized phenomenon; white matter ferritin iron levels were unchanged in patients with AD ($P = .50$).

Conclusions: The data replicate and extend prior results and suggest that basal ganglia ferritin iron levels are increased in AD. Prospective studies are needed to evaluate whether premorbid iron levels are increased in individuals who develop AD.

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The incidence rate of Alzheimer disease (AD) increases dramatically with age.1 Oxidative damage of the human brain also has been shown to be strongly age-related2,3 and has been implicated in the pathophysiology of AD.2,4-11 Tissue iron can promote oxidative damage by catalyzing free-radical reactions, resulting in the formation of hydroxyl radical (the most reactive free-radical species), which denatures protein and DNA and initiates lipid peroxidation.12 As with oxidative damage, brain iron levels increase with age.13-16 Studies of bulk brain iron have reported increased iron levels in AD brains,17-22 as have studies examining ferritin levels,19,23 a spherical protein in which upwards of 90% of tissue nonheme iron is stored.24,25 Although ferritin can sequester and store as many as 4500 iron atoms, many normal as well as pathologic processes can release iron from ferritin.12,26-29 Toxins that disrupt metabolism and result in increased lactate levels with decreasing pH will release iron from the ferritin stores.30 In addition, increases in reactive oxygen species may also release iron from ferritin,30 as do other changes, such as increased nitric oxide levels.31 Iron has been shown to interact with β-amyloid to promote neurotoxicity,32 and increased iron and ferritin has been associated with senile plaques and neurofibrillary tangles, the hallmarks of AD neuropathology.17,23,33-42

Magnetic resonance imaging (MRI) instruments produce images whose contrast depends on differences in the way tissues interact with the magnetic field of the MRI instrument. The iron atoms inside ferritin molecules form ferric oxyhydroxide particles, which profoundly affect the magnetic field, thus shortening the transverse relaxation time (T2). Magnetic resonance imaging images that are sensitive to this T2 effect (referred to as T2-weighted images) become darker in regions with high iron content (Figure 1). However, the possibility of using simple T2 measures to investigate tissue iron levels has
SUBJECTS AND METHODS

SUBJECTS

Initially, 49 subjects with AD participated in the MRI procedure after informed consent was obtained from subject and/or guardian. All participants were originally recruited from clinics associated with 2 metropolitan AD centers. Patients in these clinics undergo complete clinical assessment that includes review of clinical history, medical problems, medication use, comprehensive psychiatric and cognitive examination, blood chemistry analysis, and structural neuroimaging. All participants met National Institute of Neurological and Communicative Disorders and Stroke–Alzheimer’s Disease and Related Disorders Association (NINCDS-ADRDA) diagnostic criteria for probable or possible Alzheimer disease, as diagnosed by center-affiliated physicians. Participants were also required to speak English and have none of the following: ferromagnetic devices or implants, unstable medical illnesses or neurologic disorders that could cause cognitive deficits, history of substance use disorder within the past 2 years or prior use likely to lead to central nervous system impairment, and history of traumatic brain injury resulting in neurologic sequelae.

Eighteen of these 49 subjects were excluded from further analysis for the following reasons: 13 had untratable MRI data caused by motion or technical image artifact; 2 had a history of severe head trauma (with loss of consciousness for longer than 15 minutes) and/or seizures; 2 had a cortical infarct or a pattern consistent with multi-infarct dementia on MRI; and 1 was an outlier on the basis of age (95 years, more than 2 SDs higher than the mean age for both the AD and control groups).

The final AD population (n = 31) ranged in age from 59 to 85 years (mean, 75.6 years [SD, 6.7 years]) and consisted of 14 men and 17 women (26 whites [84%], 2 African Americans [6%], 2 Hispanics [6%], and 1 Asian [3%]); most (87% [n = 27]) were taking medications or vitamin supplements (18 [58%], psychoactive medications [eg, cholinesterase inhibitors, antidepressants]; 16 [52%], vitamin supplements [eg, E, C]; and 21 [68%], somatic medications for chronic but stable medical problems [eg, cardiovascular, arthritis]). Of the 31 subjects with AD, 27 met the NINCDS-ADRDA criteria for probable and 4 for possible AD, and all had mild to severe AD, with Mini–Mental State Examination scores ranging between 0 and 26 (mean, 17.3 [SD, 7.39]), an age of symptom onset ranging from 33.5 to 81 years (mean, 70.1 years [SD, 6.6 years]), and a length of illness (based on caregiver and/or medical record report of when symptoms were first noticed) ranging between 2 and 10 years (mean, 5.4 years [SD, 2.3 years]).

Seventy-six adult volunteers were recruited from the community and hospital staff and participated in the study as control subjects. Control subjects were excluded if they had a history of central nervous system disorder or head trauma resulting in loss of consciousness for longer than 15 minutes or if there was a family history of AD or another neurodegenerative disorder. Eight subjects were excluded from further analysis: 7 had untratable MRI data caused by technical image artifact and 1 had basal ganglia lesions on MRI examination. The final control population (n = 68) ranged in age from 59 to 82 years (mean, 68.8 years [SD, 5.3 years]) and consisted of 36 men and 32 women (59 whites [87%], 8 African Americans [12%], and 1 Asian [1%]). All control subjects denied problems with memory and scored between 27 and 30 (mean, 28.4 [SD, 0.92]) on the Mini–Mental State Examination.

MRI PROTOCOL

The methods have been described in detail elsewhere and will only be summarized herein, since the principal difference consisted of the use of a new set of MRI instruments. All subjects were scanned using the same 2 MRI instruments (1.5 T and 0.5 T; Picker Instruments, Cleveland, Ohio), and the 2 scans were done within 1 hour of each other using the same imaging protocol.

Two pilot sequences were obtained to specify the location and spatial orientation of the head and the position of the axial image acquisition grid. A coronal pilot spin–echo image (repetition time [TR], 100 milliseconds; echo time [TE], 30 milliseconds; number of excitations, 1) of 10-mm thickness was acquired and used to align the subsequent sagittal pilot images. The middle slice of the sagittal pilot images was aligned on the coronal pilot to obtain a true midsagittal image of brain. After the sagittal pilot spin–echo images (TR, 350 milliseconds; TE, 26 milliseconds; number of excitations, 2) of 5-mm thickness were acquired, the midsagittal image was used to position the axial–image acquisition grid. The axial–image acquisition sequence acquired interleaved contiguous slices using a Carr Purcell Meiboom Gill dual–spin–echo sequence (TR, 2500 milliseconds; TE, 20 and 90 milliseconds; number of excitations, 2; slice thickness, 3 mm; gradient steps, 192; field of view, 25 cm).

The coronal and sagittal pilot scans were used to determine the alignment and accuracy of head repositioning in the second MRI instrument. To consistently position the actual image slices identically within the brain and thus sample the same volume of tissue, the axial–slice select grid

been unrealized because T2 shortening is not a specific marker of tissue iron, as many tissue characteristics can affect this measure. For example, in addition to iron, the increased viscosity/density of the tightly packed myelinated axons of white matter can also shorten T2 (Figure 1). Thus, white matter has a T2 similar to cau-

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was adjusted so that the anterior commissure was contained within the same slice in both high and low field-strength instruments. For increased consistency all subsequent measures were referenced to this slice.15

**IMAGE ANALYSIS**

The T2 was calculated for each voxel by an automated algorithm from the 2 signal intensities (TEs, 20 and 90 milliseconds) of the dual-echo sequence to produce gray scale–encoded T2 maps of the brain.15

The T2 measures were extracted using an Apple Macintosh configured image analysis workstation (Apple Computer Inc, Cupertino, Calif). A single rater who was blind to clinical information obtained all measurements. The image analysis software permitted the rater to delineate the region of interest using a mouse. The contours of the entire cross-sectional area of head of the caudate, putamen, and globus pallidus and a sample of frontal lobe white matter were drawn manually by the rater using the gray and white matter contrast of the early echo (TE, 20 milliseconds) images and were then transferred onto the T2 maps.

To obtain T2 measures of homogeneous brain tissue, all pixels with T2 values that fell above the right-side inflection point on the histogram distribution of the region of interest were eliminated. This minimized the influence of voxels containing small, partial volumes of cerebrospinal fluid, which can markedly increase the T2 of the voxel. Thus, the final measure was the average T2 for the remaining homogeneous region of brain tissue.15

The T2 data for each of the 4 regions of interest were obtained from contiguous pairs of slices. The slice containing the anterior commissure and the slice immediately superior to it were used to obtain the putamen and globus pallidus T2 data. The third and fourth slices above the anterior commissure were used to obtain the T2 data for the caudate nucleus and the second and third slices superior to the orbitofrontal gray matter were used to obtain the frontal lobe white matter data. The relaxation rate (R2) was calculated as the reciprocal of T2 (1/T2). The T2 values for the 2 slices from both hemispheres were the final measures used in the subsequent analyses. The FDRI measure was calculated as the difference in R2 (high-field R2−low-field R2). Test-retest reliability for FDRI measures was very high, with intraclass correlation coefficients ranging from 0.88 to 0.99 (P<.002).15

**STATISTICAL METHODS**

As a check on the validity of the FDRI method, mean FDRI measures from the 68 control subjects in the 4 regions (Table) were correlated with postmortem iron levels in normal adults published by Hallgren and Sourander13 (Figure 2). In their study, Hallgren and Sourander13 measured non-heme iron content histologically (milligrams of iron per 100 g of fresh tissue) in human brain specimens of normal individuals aged 30 to 100 years, thus providing reference values for the various brain regions (mean±SD: caudate, 9.28 ± 2.14 [n = 58]; globus pallidus, 21.3 ± 3.49 [n = 55]; putamen, 13.32 ± 3.43 [n = 56]; and frontal white matter, 4.24 ± 0.88 [n = 59]) used to validate in vivo methods of measuring iron levels.15,16,50,54-61 Although Hallgren and Sourander13 did not report the mean age of their sample, the age range of our control subject sample (59-82 years) is within the age range of their sample (30-100 years).

An analysis of covariance design with diagnosis (AD vs controls) as the independent variable and age and sex as covariates was used to determine whether subjects with AD had increased iron in the basal ganglia. Separate analyses of covariance were computed in each region. The specificity of the results was checked by performing an additional analysis on FDRI in the white matter, where differences in iron content between AD and control subjects were not expected based on our pilot data.50 Age and sex were included as covariates in all analyses because of the known effects of age on brain ferritin iron levels15 and the possible effects of sex on tissue iron status.62 even though simple t tests (uncorrected for age and sex) result in the same significant findings as the analysis of covariance results we report. Preliminary analyses indicated that the interactions of diagnosis with age and sex were not significant, so these terms were dropped from the final models. In addition, an analysis controlling for structure volume was done to investigate possible effects of atrophy on FDRI.

Finally, separate analyses of high- and low-field R2 were done in each of the basal ganglia regions and white matter using the same analysis of covariance design. These analyses of the separate field strengths were done to evaluate whether they added diagnostically relevant information in the white matter region, as suggested in our prior report.59 Whereas the FDRI is a relatively specific measure of brain ferritin iron, high- and low-field measures of R2 are strongly influenced by other variables, such as MR visible water.

To evaluate whether illness length affected the results, the AD sample was segregated into 2 groups according to illness duration: less than 5 years (n = 15) and 5 years or more (n = 16). These 2 groups were compared by t tests for differences in age and FDRI and by χ2 test for differences in sex.

All statistical tests were 2-tailed, and α = .05 a priori.
that iron levels may be increased. In the current study compared with control subjects. This observation was in the basal ganglia of patients with AD are increased when a small sample of men suggested that ferritin iron levels in the brain. A prior pilot FDRI study performed on individuals are highly correlated with their postmortem data (Bonferroni).

Using a nonhuman primate model of hepatic hemosiderosis, a very high correlation \( r = 0.94 \) between liver tissue iron levels and field-dependent changes in \( R_2 \) was demonstrated. Finally, Hallgren and Sourander published the mean iron concentration of various brain structures of a postmortem sample of normal older individuals between the ages of 30 and 100 years (in whom age-related increases in iron content are reduced and seem to eventually plateau compared with marked age-related increases in iron content seen in younger populations). Their data has served as a reference used to compare and validate in vivo measures of brain iron, and FDRI measures of normal older individuals are highly correlated with their postmortem data (Figure 2). Thus, the FDRI measure is a specific, albeit indirect, measure of tissue ferritin iron.

The basal ganglia contain the highest levels of iron in the brain. A prior pilot FDRI study performed on a small sample of men suggested that ferritin iron levels in the basal ganglia of patients with AD are increased when compared with control subjects. This observation was in agreement with postmortem data indicating that basal ganglia iron homeostasis may be disrupted in AD and that iron levels may be increased. In the current study the FDRI method was used to evaluate brain ferritin iron in a new sample of male and female patients with AD and control subjects.

The AD and control groups differed in age (AD: mean, 75.6 years [SD, 6.7 years]; controls: mean, 68.8 years [SD, 5.3 years]) \( (t_{67} = 5.42; P < .001) \) but did not differ in sex \( \left( \chi^2 = 0.516; P = .47 \right) \) or race \( \left( \chi^2 = 5.33; P = .15 \right) \). In this older age range, correlations between age and FDRI did not reach significance for either the control or AD groups in any of the 4 structures \( (P > .25) \).

The relationship among FDRI measures of ferritin iron in caudate, putamen, globus pallidus, and white matter of our control group and published postmortem measures of iron in the same structures of normal individuals was virtually linear (Figure 2) \( (r^2 = 0.997; P = .002) \).

For the 3 basal ganglia regions, FDRI was significantly different between AD and control subjects in the caudate and putamen (Table). These differences remain significant even after experimentwise adjustment of significance levels (Bonferroni correction). Controlling for structure volume did not alter the interpretation of either result \( (P = .02 \text{ for caudate and } P = .009 \text{ for putamen}) \).

Summary statistics for \( R_2 \) in the 4 brain regions of the AD and control groups are displayed separately at each of 2 field strengths (0.5 T and 1.5 T) (Table). The \( R_2 \) data show that the high field-strength (1.5-T) instrument, which is more affected by ferritin iron than the low field-strength (0.5-T) instrument, detected differences between AD and control subjects in the caudate and putamen. However, unlike the FDRI, which is a specific measure of ferritin iron, the high field-strength \( R_2 \) differences only reached conventional levels of significance \( (P < .05, \text{ not adjusted for multiple tests [Bonferroni]}) \).

In addition, the \( R_2 \) analyses revealed highly significant differences between AD and control subjects in the white matter \( R_2 \) at both high and low field strength (Table). The difference remains significant even after Bonferroni correction.

Illness duration did not affect the results; when the AD subjects were separated into 2 groups according to illness duration, the 2 groups did not differ significantly \( (P > .30) \) in age, sex, or FDRI in any of the 4 regions of interest.
The data demonstrate significantly increased ferritin iron levels as measured by the FDRI method in the caudate and putamen of patients with AD. The data suggest that the increase in ferritin iron is specific to the caudate and putamen, since white matter ferritin iron did not differ between patients with AD and control subjects. This result is consistent with postmortem data suggesting that basal ganglia iron homeostasis may be disrupted in AD as indicated by decreased transferrin-iron ratios, increased ceruloplasmin levels (a protein that can be increased by increased iron levels), and absence of the expected age-related increase in levels of ferritin subunits in AD basal ganglia. The report of no change in ferritin levels with aging in AD suggests that the increase in FDRI we observed could be interpreted as an elevation in iron content per ferritin molecule in the basal ganglia of patients with AD. This possibility is supported by one autopsy study suggesting that ferritin from AD brains contains more iron per ferritin molecule than ferritin from normal brains.

The significantly decreased white matter $R_2$ observed in the AD group with both high- and low-field instruments (Table) could be caused by either increased MR visible water or decreased iron. The latter possibility is refuted by the FDRI data (Table). Therefore, the significantly decreased white matter $R_2$ observed in the AD group is consistent with earlier MRI reports suggesting processes that increase MR visible water, such as myelin loss or tissue water increase, in the AD brain.

As shown in the Table, low-field $R_2$ seems to be a relatively pure measure of this second field-strength-independent process that decreases $R_2$ in the white matter of AD. The low-field instrument also shows no evidence of increased $R_2$ values in the caudate or globus pallidus of patients with AD since it is minimally affected by ferritin. Thus, the low-field $R_2$ measure provides a basis for "correction" of the high-field $R_2$ measure, subtracting variance associated with field-strength-independent effects on $R_2$. Conversely, although the high-field $R_2$ measure is affected by ferritin iron and can detect differences in basal ganglia ferritin iron levels between the control and AD groups (Table), better discrimination is obtained with the FDRI method since the variance associated with field-strength-independent effects is eliminated (Table).

Important limitations of this study need to be considered when interpreting the data. The cross-sectional design could obscure some of the AD-vs-control differences in FDRI and $R_2$. This could happen if brain iron levels are associated with mortality and/or accelerated debilitation that cause patients to be excluded from, unable to participate in, or unable to complete the study. In addition, in a cross-sectional study, only limited inferences can be made about the causal relationship between increased iron levels and AD pathologic findings. It was not feasible to examine neocortical structures in this study, despite their relevance to AD. Accurate FDRI measurements require that the same tissue be identified and measured in the 2 different MRI scans. At present, this is technically difficult since the cortex is very thin and convoluted. Finally, the AD and control groups differed in mean age. However, in this age range (59 years and older), age and FDRI were not significantly correlated for either the control or AD group, and in addition, age was controlled for statistically by including it as a covariate in the analyses of covariance.

We attempted to evaluate whether the increased ferritin iron was simply a marker for disease-related cell loss that secondarily results in higher iron concentrations or whether the iron contributes to the pathogenesis of the AD disease processes and aging, as others have suggested. Differences in FDRI could not be accounted for solely by a reduction in the volume of the structures. Therefore, the observed changes in this study cannot be attributed solely to structure atrophy. Similarly, the length of illness seems to have no impact on basal ganglia ferritin iron levels, suggesting that the ferritin iron increases in AD are not caused by the illness itself and may be interpreted as a risk factor for AD.

An association between high iron levels and central nervous system damage has been observed in a va-
variety of neurodegenerative disorders, and iron involvement in the process of oxidative damage has been suggested as a common mechanism. In addition to AD, the involvement of iron and free-radical neurotoxic processes has been postulated for a variety of age-related neuropsychiatric disorders, such as Parkinson disease, Huntington disease, and tardive dyskinesia. Since brain iron levels increase with age, the risk of iron-mediated damage may increase with age, suggesting a role for iron in age-related neurodegenerative disorders.

In vivo ferritin iron quantification could lead to improved diagnostic and prognostic criteria for AD. If a causal relationship is established between premorbid iron levels and AD through prospective in vivo MRI studies, new avenues of treatment and prevention of the disease may be possible. In addition, serial evaluations of FDRI could be used to monitor iron chelating treatments, which are currently used in patients with transfusion hemosiderosis and have been tentatively shown to be effective in the treatment of some patients with AD.

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