Iron and Non-Iron-Related Characteristics of Multiple Sclerosis and Neuromyelitis Optica Lesions at 7T MRI


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ABSTRACT

BACKGROUND AND PURPOSE: Characterization of iron deposition associated with demyelinating lesions of multiple sclerosis and neuromyelitis optica has not been well studied. Our aim was to investigate the potential of ultra-high-field MR imaging to distinguish MS from neuromyelitis optica and to characterize tissue injury associated with iron pathology within lesions.

MATERIALS AND METHODS: Twenty-one patients with MS and 21 patients with neuromyelitis optica underwent 7T high-resolution 2D-gradient-echo-T2* and 3D-susceptibility-weighted imaging. An in-house-developed algorithm was used to reconstruct quantitative susceptibility mapping from SWI. Lesions were classified as “iron-laden” if they demonstrated hypointensity on gradient-echo-T2*-weighted images and/or SWI and hyperintensity on quantitative susceptibility mapping. Lesions were considered “non-iron-laden” if they were hyperintense on gradient-echo-T2* and iso- or hypointense on quantitative susceptibility mapping.

RESULTS: Of 21 patients with MS, 19 (90.5%) demonstrated at least 1 quantitative susceptibility mapping–hyperintense lesion, and 11/21 (52.4%) had iron-laden lesions. No quantitative susceptibility mapping–hyperintense or iron-laden lesions were observed in any patients with neuromyelitis optica. Iron-laden and non-iron-laden lesions could each be further characterized into 2 distinct patterns based on lesion signal and morphology on gradient-echo-T2*/SWI and quantitative susceptibility mapping. In MS, most lesions (n = 262, 75.9% of all lesions) were hyperintense on gradient-echo-T2* and iso- or hypointense on quantitative susceptibility mapping (pattern A), while a small minority (n = 26, 7.5% of all lesions) were hyperintense on both gradient-echo-T2* and quantitative susceptibility mapping (pattern B). Iron-laden lesions (n = 57, 16.5% of all lesions) were further classified as nodular (n = 22, 6.4%, pattern C) or ringlike (n = 35, 10.1%, pattern D).

CONCLUSIONS: Ultra-high-field MR imaging may be useful in distinguishing MS from neuromyelitis optica. Different patterns related to iron and noniron pathology may provide in vivo insight into the pathophysiology of lesions in MS.

ABBREVIATIONS: GRE = gradient-echo; NMO = neuromyelitis optica; ppb = parts per billion; QSM = quantitative susceptibility mapping; R2* = transverse relaxation rate

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ORIGINAL RESEARCH

ADULT BRAIN

MULTIPLE SCLEROSIS AND NEUROMYELITIS OPTICA (NMO) IS A DISTINCT DISORDER OF THE CNS WITH DIFFERENT PATHOPHYSIOLOGY AND CLINICAL MANIFESTATIONS THAN MULTIPLE SCLEROSIS (MS). Distinguishing these 2 conditions can be challenging in view of the overlap in clinical manifestations and findings on conventional neuroimaging.2 Ultra-high-field imaging, by virtue of increased SNR, increased spatial resolution, and markedly improved venous and iron contrast within lesions,3 has the potential to shed light on the underlying pathophysiology of MS and NMO and help distinguish these 2 conditions. 7T MR imaging studies4,5 have shown that 60%–80% of MS lesions are traversed by a central venule, while only a small minority of NMO lesions contain a central venule.2 Differentiation of MS from NMO before the commencement of treatment may be challenging in view of the overlap in clinical manifestations and findings on conventional neuroimaging.5 However, ultra-high-field imaging, by virtue of increased SNR, increased spatial resolution, and markedly improved venous and iron contrast within lesions,4 has the potential to shed light on the underlying pathophysiology of MS and NMO and help distinguish these 2 conditions. 7T MR imaging studies4,5 have shown that 60%–80% of MS lesions are traversed by a central venule, while only a small minority of NMO lesions contain a central venule.2 Differentiation of MS from NMO before the commencement of treatment may be challenging in view of the overlap in clinical manifestations and findings on conventional neuroimaging.5 However, ultra-high-field imaging, by virtue of increased SNR, increased spatial resolution, and markedly improved venous and iron contrast within lesions,4 has the potential to shed light on the underlying pathophysiology of MS and NMO and help distinguish these 2 conditions. 7T MR imaging studies4,5 have shown that 60%–80% of MS lesions are traversed by a central venule, while only a small minority of NMO lesions contain a central venule.2 Differentiation of MS from NMO before the commencement of treatment may be challenging in view of the overlap in clinical manifestations and findings on conventional neuroimaging.5 However, ultra-high-field imaging, by virtue of increased SNR, increased spatial resolution, and markedly improved venous and iron contrast within lesions,4 has the potential to shed light on the underlying pathophysiology of MS and NMO and help distinguish these 2 conditions. 7T MR imaging studies4,5 have shown that 60%–80% of MS lesions are traversed by a central venule, while only a small minority of NMO lesions contain a central venule.2 Differentiation of MS from NMO before the commencement of treatment may be challenging in view of the overlap in clinical manifestations and findings on conventional neuroimaging.5 However, ultra-high-field imaging, by virtue of increased SNR, increased spatial resolution, and markedly improved venous and iron contrast within lesions,4 has the potential to shed light on the underlying pathophysiology of MS and NMO and help distinguish these 2 conditions. 7T MR imaging studies4,5 have shown that 60%–80% of MS lesions are traversed by a central venule, while only a small minority of NMO lesions contain a central
MATERIALS AND METHODS

Subjects

This study was conducted at 2 academic MS referral centers: New York University Medical Center, New York, and Charité University, Berlin. Both sites received approval from local institutional review boards. Written informed consent was obtained from all patients before study entry. Inclusion criteria were a diagnosis of NMO spectrum disorders (International Panel for NMO Diagnosis Criteria) or definite MS (McDonald Criteria). Twenty-one patients with MS (mean age, 47.6 ± 14.2 years; all women, mean disease duration, 8.4 ± 6.7 years; range, 1–26.6 years) were enrolled in this study. All patients were NMO Ab seropositive by immunohistochemical or by enzyme-linked immunosorbent assays.

Twenty-one patients with MS were enrolled (mean age, 47.1 ± 10.3 years; 6 men/15 women; disease duration, 11.5 ± 5.9 years; range, 4–25 years). This group included patients with relapsing-remitting (n = 19) and secondary-progressive (n = 2) MS. There was no significant difference in mean age and mean disease duration between MS and NMO groups (P > .05).

Ultra-High-Field MR Imaging

All patients underwent ultra-high-field MR imaging by using identical whole-body 7T human MR imaging systems (Magnetom; Siemens, Erlangen, Germany) equipped with a 24-channel phased array coil (Nova Medical, Wilmington, Massachusetts). The imaging protocol included high-resolution axial 2D-gradient-echo (GRE) T2*-weighted imaging, high-resolution axial 3D-SWI, FLAIR, and sagittal T1-weighted 3D-MPRAGE sequences. Only supratentorial brain regions were covered while acquiring 2D-GRE-T2* and 3D-SWI to avoid susceptibility artifacts from air-tissue interfaces. None of the patients received intravenous contrast agent. The acquisition parameters were the following: for GRE-T2*-weighted imaging: TR/TE = 580/25 ms, flip angle = 35°, section thickness = 2 mm, FOV = 240 × 240 mm², voxel size = 0.2 × 0.2 × 2 mm³; for sagittal 3D sampling perfection with application-optimized contrasts by using different flip angle evolution sequence (SPACE; Siemens) FLAIR: TR/TE/ TI = 8000/380/2100 ms, isotropic voxel size = 1.0 × 1.0 × 1.0 mm³; and for sagittal T1-weighted 3D-MPRAGE: TR/TE/TI = 2000/2.92/1100 ms, isotropic voxel size = 1.0 × 1.0 × 1.0 mm³.

High-resolution flow-compensated 3D-SWI was acquired with the following parameters: TR/TE = 27/18 ms, flip angle = 18°, section thickness = 2 mm, FOV = 240 × 240 mm², base resolution = 1024, voxel size = 0.2 × 0.2 × 2 mm³, bandwidth = 110 Hz/px, acquisition time = 7 minutes and 49 seconds, and integrated parallel imaging technique factor = 2. Equivalent imaging parameters were used at both sites for all the sequences; however, at the German site, a voxel size of 0.5 × 0.5 × 2 mm³ was used for the 3D-SWI sequence.

Data Postprocessing

The source magnitude and phase images from each SWI scan were obtained and used to generate SWI venography. All phase images were reconstructed and corrected for field inhomogeneities with a Hamming high-pass filter (96 × 96) by using Signal Processing In NMR software (SPIN; MR Imaging Institute for Biomedical Research, Detroit, Michigan). The original magnitude image was multiplied by the phase mask 4 times to enhance the visibility of lesion and venous structures. Finally, SWI venograms were created by performing minimum intensity projection over 2 contiguous sections.

Susceptibility-weighted imaging and the mapping algorithm developed by Haacke et al. were used to reconstruct QSM maps from high-resolution 3D-SWI data. The postprocessing involved skull stripping to remove the artifacts caused by skull and brain tissue interface by using the FSL Brain Extraction Tool (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/BET), followed by phase unwrapping by using a Laplacian operator. To remove background field inhomogeneity, we applied a variable high-pass filter of 32 pixels and, finally, performed inverse filtering to generate QSM maps.

Data Analysis

All MR images were analyzed by using ImageJ software (National Institutes of Health, Bethesda, Maryland). All MS and NMO lesions were analyzed side by side on axial GRE-T2*, 3D-SWI, and QSM images. The following morphologic imaging characteristics were recorded for each lesion: 1) the largest cross-sectional diameter, 2) the presence of 1 or multiple central intraleSIONAL veins, 3) differential signal intensity within the lesions, and 4) the presence of a peripheral rim.

The lesion signals on GRE-T2*, SWI, and QSM and their putative correlations with underlying pathology are summarized in
The mean cross-sectional diameter for MS lesions (5.44 ± 2.66 mm) was significantly larger than that of NMO lesions (3.19 ± 1.12 mm, P < .001). While MS lesions had variable shapes, most of the NMO lesions were round. Of 345 MS lesions, 227 (65.8%) were traversed by intralesional central venules, while only 8/132 (6.1%) NMO lesions showed this feature. All patients with MS had ≥1 lesion traversed by a central venules. On the other hand, only 4 patients with NMO had ≥1 lesion traversed by a central venules. The presence of at least 1 lesion with a central venule distinguished MS from NMO with a sensitivity of 100% (95% CI, 83.9%–100%) and a specificity of 71.4% (95% CI, 47.8%–88.7%).

MS lesions varied in signal intensity on GRE-T2*, SWI, and QSM. In the MS cohort, 19/21 (90.5%) patients had at least 1 hyperintense lesion on QSM, and this feature distinguished patients with MS from those with NMO with a sensitivity of 90.5% (95% CI, 69.6%–98.8%) and a specificity of 100% (95% CI, 83.9%–100%). Moreover, 11/21 (52.4%) patients had iron-laden lesions (see the “Data Analysis” section for a definition). In contrast, all NMO lesions demonstrated hyperintense signal relative to surrounding brain parenchyma on GRE-T2* weighted images and were isointense (inconspicuous) on QSM. Thus, none of the patients with NMO had any iron-rich lesions (Fig 1). The presence of at least 1 iron-laden lesion characteristic distinguished MS from NMO with a sensitivity of 52.4% (95% CI, 29.8%–74.3%) and a specificity of 100% (95% CI, 83.9%–100%).

Characterization of Different Lesion Patterns in MS

Examples of 4 morphologically distinct lesion patterns seen in MS are shown in Figs 2 and 3. Signal intensity on GRE-T2*, SWI, and QSM for different lesion patterns in MS is presented in Table 2. Most of the MS lesions (n = 262, 75.9%) were pattern A (all were hyperintense on GRE-T2*-weighted images and inconspicuous on QSM). On SWI, only 53/262 of these lesions demonstrated hyperintensity. Of 262 pattern A lesions, 148 (56.4%) were traversed by a central venule. A small number of lesions (n = 26, 7.5%) were pattern B (hyperintensity on both GRE-T2*-weighted images and QSM). On SWI, only 11/26 of these lesions demonstrated hyperintensity. Of the 26 pattern B lesions, 22 (84%) were traversed by a central venule.

Some lesions (n = 57, 16.5% of all MS lesions) demonstrated hyperintensity on QSM but were inconspicuous on GRE-T2*. Of the 132 NMO patients, only 8/132 (6.1%) showed hyperintensity on QSM. On the other hand, 26 (7.8%) of these lesions also showed hyperintensity on GRE-T2*. A small number of lesions (n = 22, 6.1%) were pattern C (hyperintense on GRE-T2* and isointense or hyperintense on QSM). These lesions were further classified as nodular (pattern C) or exhibiting a peripheral rim (pattern D). We computed QSM values from all lesions that demonstrated hyperintensity on QSM by manually drawing ROIs. Because the use of a high-pass filter while reconstructing QSM maps may reduce the effective iron content from different tissue compartments, we believe this process might have resulted in underestimation of the QSM values computed from iron- and non-iron-laden lesions. To correct for the QSM values, we used a simulation algorithm24 to obtain a scaling factor based on the size of the lesions. This size-dependent scaling factor was multiplied by the original QSM values to obtain corrected QSM values for each lesion.

Table 1: Proposed histopathologic interpretation based on signal-intensity changes on MR images

<table>
<thead>
<tr>
<th>Tissue Content</th>
<th>Susceptibility Effect</th>
<th>Signal Intensity on GRE-T2*</th>
<th>Signal Intensity on SWI</th>
<th>Signal Intensity on QSM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium</td>
<td>Diamagnetism</td>
<td>Hypointense</td>
<td>Hypointense</td>
<td>Hypointense</td>
</tr>
<tr>
<td>Myelin</td>
<td>Diamagnetism</td>
<td>Hypointense</td>
<td>Isointense</td>
<td>Isointense</td>
</tr>
<tr>
<td>Variable degree of micronecrosis, edema, gliosis, demyelination, and macromolecules</td>
<td>Diamagnetism</td>
<td>Hyperintense</td>
<td>Isointense or hyperintense</td>
<td>Isointense</td>
</tr>
<tr>
<td>Extensive degree of demyelination</td>
<td>Loss of diamagnetism (paramagnetism)</td>
<td>Hyperintense</td>
<td>Isointense or hyperintense</td>
<td>Hyperintense</td>
</tr>
<tr>
<td>Iron</td>
<td>Paramagnetism</td>
<td>Hypointense</td>
<td>Hypointense</td>
<td>Hyperintense</td>
</tr>
</tbody>
</table>

**Statistical Analysis**

A χ² test was performed to look for differences in proportions of lesions with central venules in patients with MS and NMO. Receiver operating characteristic analyses were performed to estimate the sensitivity and specificity of selected subject-level conditions (at least 1 lesion with a central venule, at least 1 iron-laden lesion, at least 1 QSM hyperintense lesion) as criteria for classifying patients as testing positive for MS. Clopper-Pearson confidence intervals were derived for the sensitivity and specificity associated with each condition.

All lesions providing a QSM value were classified as having a central venule. Mixed-model analysis of variance was used to compare iron-laden from non-iron-laden lesions in terms of QSM while accounting for the correlation among QSM values derived for lesions in the same patient. The covariance structure was modeled by assuming QSM values to be independent when acquired from different patients and symmetrically correlated when acquired from lesions within the same patient, with the strength of correlation dependent on whether lesions were of the same type (eg, lesions were both classified as iron-laden). A probability value < .05 was considered significant. All data analysis was performed by using SPSS for Windows, Version 15.0 (IBM, Armonk, New York).

**RESULTS**

**Distinguishing Patients with MS from Those with NMO**

A total of 345 MS and 132 NMO discrete lesions were observed in supratentorial brain regions of 21 patients with MS and 21 with NMO. The mean cross-sectional diameter for MS lesions (5.44 ± 2.66 mm) was significantly larger than that of NMO lesions (3.19 ± 1.12 mm, P < .001). While MS lesions had variable shapes, most of the NMO lesions were round. Of 345 MS lesions, 227 (65.8%) were traversed by intralesional central venules, while only 8/132 (6.1%) NMO lesions showed this feature. All patients with MS had ≥1 lesion traversed by a central venule. On the other hand, only 4 patients with NMO had ≥1 lesion traversed by a central venule. The presence of at least 1 lesion with a central venule distinguished MS from NMO with a sensitivity of 100% (95% CI, 83.9%–100%) and a specificity of 71.4% (95% CI, 47.8%–88.7%).

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hypointensity on GRE-T2*-weighted images and/or SWI, but hyperintensity on QSM; these lesions were presumed to be iron-laden. Seventeen of 57 iron-laden MS lesions demonstrated hyperintensity on GRE-T2*, and 40/57 were hypointense on GRE-T2* images. All except 1 (56/57) demonstrated hypointensity on SWI. All of these iron-laden QSM hyperintense lesions were traversed by a central venule (57/57). Furthermore, of 288 non-iron-laden MS lesions, 240 (83.3%) had ill-defined and faint margins generally, while only 12 of 57 (21%) iron-laden MS lesions had ill-defined margins (Fig 4).

The iron-laden lesions had 2 geometrically distinct susceptibility patterns: nodular if they had solid signal intensity on QSM (n = 22, 6.4%; pattern C) or ringlike if they had a distinct peripheral rim on QSM (n = 35, 10.1%; pattern D). However, all of these iron-laden lesions demonstrated hypointensity on SWI. The mean QSM was significantly higher for iron-laden lesions than for non-iron-laden lesions (P = .027; Fig 5A). The least squares mean standard error of the mean of QSM, adjusted for within-subject correlations, was 38.73 ± 4.81 parts per billion (ppb) for iron-laden lesions and 26.36 ± 2.2 ppb for lesions that were not iron-laden. Receiver operating characteristic analysis provided a threshold QSM value of 39.26 ppb to distinguish iron- from non-iron-laden lesions with a sensitivity of 70.2% and specificity of 57.7% (Fig 5B).

**DISCUSSION**

Early differentiation of NMO from MS is crucial for optimal management clinically. Although the recent availability of commercial testing for antibodies to Aquaporin-4 water has facilitated differentiation of NMO from MS, a correct diagnosis still remains challenging, particularly in those patients with NMO with multiple brain lesions on MR imaging. Many patients with NMO are still misdiagnosed with MS. Hence, development of newer imaging biomarkers that can enable objective separation of these 2 diseases is warranted. Our analysis of the morphologic and structural features of MS and NMO supratentorial lesions by using multicontrast 7T MR imaging helps to further differentiate the 2 conditions and provides insight into lesional pathology in vivo. We found that iron deposition within a lesion (hyperintense signal on QSM) distinguished patients with MS from those with NMO with a sensitivity of 90.5% (95% CI, 69.6%–98.8%) and a specificity of 100% (95% CI, 83.9%–100%). Analysis of the signal intensity of lesions on GRE, SWI, and QSM sequences allowed us to divide all lesions into 4 patterns, of which 2 were iron-enriched and 2 were non-iron-enriched (Table 2). A characteristic feature of MS was the...
variety of lesion types observed: All patients with MS had lesions of ≥1 type, though most lesions were of pattern A, while all NMO lesions were of 1 type only (pattern A).

Using phase imaging and transverse relaxation rate (R2*) mapping on high-field MR imaging, prior studies have assessed the iron content of MS lesions qualitatively and quantitatively. However, many confounding factors are associated with these imaging techniques, including dependence on orientation and distribution of susceptibility sources that may influence the heterogeneity of phase and R2* and thereby render these images less reliable and quantifiable. QSM, on the other hand, provides more robust and quantitative evaluation of magnetic susceptibility sources such as iron, myelin, and calcium that are present in normal and diseased brain tissues. Moreover, susceptibility is a physical quantity that is independent of imaging parameters and has the potential to distinguish and quantify different susceptible tissues. In the present study, a recently described susceptibility-weighted imaging and mapping method was used to reconstruct QSM from 3D-SWI, providing susceptibility values from different tissue compartments with high accuracy.

Our findings on 7T MR imaging that not all MS lesions have iron deposition are consistent with the earlier, lower field studies. Iron deposition may vary among individual lesions on the basis of their age and inflammatory status. We observed 4 morphologically dis-

**Pattern C**

**Pattern D**

**Figure 3.** Iron-laden MS lesions (2 geometrically distinct susceptibility patterns). In the upper rows, schematic sketches are shown for pattern C and D lesions, depicting a characteristic signal-intensity distribution on GRE-T2*-weighted images, SWI, and QSM. In the lower rows, axial GRE-T2*-weighted image presents a pair of lesions in the subcortical WM region. While one lesion is nodular hyperintense (white arrow), another lesion is nodular hypointense (black arrow). Both lesions are crisscrossed by intralesional venules and show hypointense signal on SWI but hyperintense signal on QSM, thus indicating iron deposition (pattern C). Axial GRE-T2*-weighted image reveals a lesion (white arrow) with a hypointense peripheral rim having a central venous structure in the subcortical WM region. The lesion also shows a hypointense rim on the corresponding SWI and a hyperintense rim on the corresponding QSM, suggesting a ringlike lesion with iron deposition only at the edges (pattern D).

**Table 2: Signal intensity on GRE-T2*, SWI, and QSM for different lesion patterns in MS**

<table>
<thead>
<tr>
<th>Lesion Pattern</th>
<th>Signal Intensity on GRE-T2*</th>
<th>Signal Intensity on SWI</th>
<th>Signal Intensity on QSM</th>
</tr>
</thead>
<tbody>
<tr>
<td>A (n = 262)</td>
<td>Hyperintense</td>
<td>Isointense</td>
<td>Isointense</td>
</tr>
<tr>
<td>B (n = 26)</td>
<td>Hyperintense</td>
<td>Isointense or hypointense</td>
<td>Hyperintense</td>
</tr>
<tr>
<td>C (n = 22)</td>
<td>Hyperintense or hypointense</td>
<td>Hyperintense or hypointense</td>
<td>Hyperintense</td>
</tr>
<tr>
<td>D (n = 35)</td>
<td>Hyperintense or hypointense</td>
<td>Hyperintense or hypointense</td>
<td>Hyperintense</td>
</tr>
</tbody>
</table>

Iron-laden MS lesions with hyperintensities on QSM were of 2 patterns: iron either deposited in the center core (nodular iron-laden lesions, pattern C) or in a ringlike fashion at the lesion edge (pattern D). Previous histochemical studies showed that iron deposits were present in a subset of chronic, demyelinating active MS lesions. The molecular pathways for iron accumulation in MS lesions are still not fully understood; however, several possible biologic mechanisms, such as iron-rich oligodendrocyte debris, iron-sequestered microglia or macrophages, and products of local microhemorrhages following venule wall damage may contribute to iron deposition. On 3T MR imaging, Chen et al. observed lower QSM values from acute enhancing lesions (small susceptibilities) and increased QSM values from nonenhancing lesions (high susceptibilities). This observation suggests that lesion susceptibility measured by QSM is a useful biomarker for monitoring MS disease activities. Perhaps, the preponderance of non-iron-containing lesions in MS in our study is because they were imaged long after their formative stage.

Most iron-laden lesions in our study were ringlike. Histopathologically, it has been observed that maximum accumulation of iron occurs at the edges of classic, reactive, slowly expanding chronically existing MS lesions. In these reactive lesions, demyelination and oligodendrocyte destruction occur in a zone of variable size at the lesion border. Thus, iron-containing myelin and oligodendrocytes gradually decreased from the perilesional regions toward the lesion centers. Additionally, iron-containing microglia and macrophages that are mainly located at the edge of chronic reactive lesions undergo microglial dystrophy leading to a variable degree of iron deposition within the different compartments of the MS lesions. Moreover, iron-laden MS lesions were well-circumscribed with well-defined margins while non-iron-laden lesions had poorly defined margins and were generally larger. The reason for this observation is not clearly understood; however, it might be because of accumulation of...
iron-enriched microphages or microglia cells in the iron-laden lesions or more activity associated with these lesions.

An interesting observation to emerge from our study is that the probability of finding a small venule in the center of lesions depends on the lesion pattern. All of the iron-laden MS lesions, 100% (patterns C and D), had significantly higher QSM than non-iron-laden QSM hypointense lesions (pattern B). Boxes represent the median, 25th percentile, and 75th percentile. The asterisk indicates a significant difference (P = .001). Solid circles represent the outliers. The receiver operating characteristic curve shows an area under the curve of 0.74. The receiver operating characteristic analysis (B) provides a sensitivity of 70.2% and a specificity of 57.7% in distinguishing these 2 types of lesions at a threshold QSM value of 30.26 ppb.

Another interesting observation was that only 70% of the iron-laden lesions demonstrated hypointense signal on GRE-T2*-weighted images, whereas 98% of the lesions showed hypointensity on SWI, despite similar section thickness and in-plane resolution for both of these images. In accordance with previous studies, our observation suggests that SWI is more sensitive to susceptibility effects than GRE-T2*-weighted images, probably because SWI combines information both from phase and magnitude images to ascertain the local susceptibility changes among neighboring tissues, whereas susceptibility contrast on GRE-T2* is mainly dependent on a combination of spin-spin relaxation (T2) and magnetic field inhomogeneity. We believe that inclusion of phase information renders SWI more sensitive to susceptibility effects than GRE-T2* images.

There were some limitations to the current study. Most of the patients with MS had a diagnosis of relapsing-remitting MS. To
obtain a more comprehensive understanding of the evolution of MS lesions by using multicontrast imaging, future studies would need to include patients with different types of MS (clinically isolated syndrome, secondary- and primary-progressive MS). Our patients with NMO had similar kinds of “nonspecific” subcortical lesions. A number of other kinds of NMO-specific lesions have been described but were not seen in our series. Another limitation of the current study was that R2* or T2* mapping was not performed; these sequences may provide additional quantitative information for the characterization of NMO and MS lesions.

CONCLUSIONS
Four morphologically different patterns were observed for MS lesions, while NMO lesions exhibited only 1 pattern. Approximately half of patients with MS (52.4%) had at least 1 iron-laden lesion, but none of the patients with NMO had iron-enriched lesions. Our study suggests that QSM, combined with other imaging sequences at 7T, helps further differentiate MS from NMO and provides insight into lesional pathogenesis and iron metabolism in these 2 autoimmune disorders of the central nervous system.

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REFERENCES


