Quantitative Susceptibility Mapping and R2* Measured Changes during White Matter Lesion Development in Multiple Sclerosis: Myelin Breakdown, Myelin Debris Degradation and Removal, and Iron Accumulation


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ABSTRACT

BACKGROUND AND PURPOSE: Quantitative susceptibility mapping and R2* are sensitive to myelin and iron changes in multiple sclerosis lesions. This study was designed to characterize lesion changes on quantitative susceptibility mapping and R2* at various gadolinium-enhancement stages.

MATERIALS AND METHODS: This study included 64 patients with MS with different enhancing patterns in white matter lesions: nodular, shell-like, nonenhancing < 1 year old, and nonenhancing 1–3 years old. These represent acute, late acute, early chronic, and late chronic lesions, respectively. Susceptibility values measured on quantitative susceptibility mapping and R2* values were compared among the 4 lesion types. Their differences were assessed with a generalized estimating equation, controlling for Expanded Disability Status Scale score, age, and disease duration.

RESULTS: We analyzed 203 lesions: 80 were nodular-enhancing, of which 77 (96.2%) were isointense on quantitative susceptibility mapping; 33 were shell-enhancing, of which 30 (90.9%) were hyperintense on quantitative susceptibility mapping; and 49 were nonenhancing lesions < 1 year old and 41 were nonenhancing lesions 1–3 years old, all of which were hyperintense on quantitative susceptibility mapping. Their relative susceptibility/R2* values were 0.5 ± 4.4 parts per billion/−5.6 ± 2.9 Hz, 10.2 ± 5.4 parts per billion/−8.0 ± 2.6 Hz, 20.2 ± 7.8 parts per billion/−3.1 ± 2.3 Hz, and 33.2 ± 8.2 parts per billion/−2.0 ± 2.6 Hz, respectively, and were significantly different (P < .005).

CONCLUSIONS: Early active MS lesions with nodular enhancement show R2* decrease but no quantitative susceptibility mapping change, reflecting myelin breakdown; late active lesions with peripheral enhancement show R2* decrease and quantitative susceptibility mapping increase in the lesion center, reflecting further degradation and removal of myelin debris; and early or late chronic nonenhancing lesions show both quantitative susceptibility mapping and R2* increase, reflecting iron accumulation.

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THE PATHOLOGIC CHANGES IN ACTIVE WHITE MATTER MS LESIONS INCLUDE INFLAMMATORY INfiltrATION WITH IMMUNE CELLS, MYELIN BREAKDOWN AND REMOVAL BY MICROGLIA/MACROPHAGES, AND IRON ACCUMULATION IN IMMUNE CELLS WITHIN THE LESION AND THE LESION PERIPHERY.1,2

MR imaging staging of white-matter MS lesions uses gadolinium (Gd) enhancement to differentiate between active (enhancing) and nonactive (nonenhancing) lesions. However, enhancement provides only a short window into inflammatory activity...
because it is preceded and outlasted by infiltration with immune cells. Enhancement on T1WI + Gd reflects BBB damage, whereas conventional MR imaging (T2WI and T1WI) sensitizing water mobility does not reflect specific microstructural and compositional changes in active lesions. In contrast, gradient-echo (GRE) imaging is sensitive to the magnetic fields induced by inhomogeneous myelin and highly paramagnetic iron, offering the potential to study microstructural and compositional changes in active MS lesions. GRE imaging data can be processed to generate quantitative susceptibility mapping (QSM) and R2* maps. QSM measures the underlying total magnetic susceptibility sources in a voxel. R2* reflects the sum of the T2 relaxation rate that depends on cellular content and the field dispersion associated with susceptibility microstructural heterogeneity multiplied by TE. Breakdown of intact myelin decreases R2* but does not change QSM signal, whereas degradation of diamagnetic myelin debris within macrophages and removal of degraded myelin increase susceptibility measured on QSM. Finally, iron accumulation increases both R2* and QSM. Therefore, by using both QSM and R2*, we would be able to differentiate between acute myelin breakdown, advanced myelin degradation and removal, and iron accumulation in MS lesions.

**MATERIALS AND METHODS**

**Patients**

The Weill Cornell Medical College institutional review board approved this MR imaging study of patients with MS from August 2011 to April 2015. The inclusion criteria were patients who 1) had undergone, within 12 months, 2 successive MRIs that included T2WI, and pre- and post-Gd T1WI, with the second MR imaging including a multi-echo GRE imaging; and 2) had new hyperintense white matter lesions on the second T2WI. A total of 65 patients met the inclusion criteria; 1 patient was excluded a posteriori because of motion artifacts on GRE images. The mean age of the 64 patients (13 men and 51 women) was 35.29 ± 8.12 years, with age ranging from 11–51 years. Patients were receiving the steroid Solu-Medrol intravenous injection (3 mg/day for 3 to 5 days). All their disease durations ranged from 0–17 years (6.0 ± 4.71 years) and the Expanded Disability Status Scale scores ranged from 0–6 (median, 1.5; interquartile range, 3).

**Imaging Protocol and Reconstruction**

MR imaging was performed on a 3T MR scanner (Signa HDxt; GE Healthcare, Milwaukee, Wisconsin) with an 8-channel head coil. The sequences for each patient were: 1) T2-weighted FSE (TR/TE = 5250/86 msec, flip angle = 90°, 3-mm-thick section at 0 intervals, 416 × 256 matrix, 24-cm FOV); 2) pre- and 3) post-Gd 3D inversion recovery-prepared T1-weighted fast echo-spoiled gradient-echo (TR/TE = 8.8/3.4 msec, flip angle = 15°, 256 × 256 matrix, 24-cm FOV, 0.45 × 0.45 × 1.2 mm³ resolution); and 4) 3D T2*-weighted spoiled multiecho GRE sequence. Imaging parameters for the multiecho GRE sequence were as follows: TR = 57 msec; number of echoes = 11; first TE = 4.3 msec; TE spacing = 4.8 msec; flip angle = 20°; bandwidth = 244 kHz; FOV = 24 cm; thickness = 2 mm; acquisition matrix = 416 × 320; acquisition voxel size = 0.5 × 0.5 × 2 mm³. The GRE sequence was performed before Gd injection, and the post-Gd T1WI was performed at 4 minutes after Gd injection.

QSM was reconstructed from the data acquired with the GRE sequence by using the morphology-enabled dipole inversion method, and R2* maps were generated by using a Gauss-Newton least-squares mono-exponential fit of the 11 echoes. QSM and R2* were reconstructed to a matrix of 512 × 512, resulting in a voxel size of 0.47 × 0.47 × 2 mm³. All images of 1 patient were registered to the QSM image by using the FMRIB Linear Image Registration Tool (FLIRT; http://www.fmrib.ox.ac.uk/).

**Lesion Analysis**

New white matter MS lesions were identified on the second MR imaging, which was performed at 0.77 ± 0.37 year after the first MR imaging. Three neuroradiologists (J.C., A.G., and G.C.-Y.C., with 18, 9, and 8 years of experience, respectively) independently...
reviewed all images of the selected patients and classified new lesions into 3 groups based on enhancement on T1WI + Gd (nodular-enhancing [solid in 2D], shell-enhancing [ringlike in 2D], and nonenhancing < 1 year old). By comparison to an MR imaging < 3 years old, lesions 1–3 years old (1.57 ± 0.46 years; error, 0.88 ± 0.34 years) from the same cohort were identified as a fourth group. The method of estimating lesion age referred to a previous study. All these lesions were independently assessed by the 3 neuroradiologists to be hyperintense or isointense on QSM images. These 3 readings were combined, with all differences resolved by majority votes.

ROIs of the lesions were placed on the lesion’s hyperintense area on T2-weighted images, but the edema around was not included. If a lesion had a hypointense rim on T2WI, we also excluded this rim (example is shown on Fig 1). ROIs as normal-appearing white matter references were drawn on the contralateral mirror site of the lesions with similar shape and size.
One neuroradiologist (Y.Z., 4 years of experience) drew the ROIs on T2WI and then overlaid them on QSM and R2* maps by using in-house ROI semiautomatic software implemented for OsiriX (http://www.osirix-viewer.com). The veins and artifacts inside lesions on QSM and R2* were carefully removed by hand.

**Statistical Analysis**

The differences in relative susceptibilities and R2* values of lesions in the 4 groups were assessed by a generalized estimating equation. This model assumes a Gaussian distribution and an exchangeable correlation structure to account for the multiple QSM and R2* measurements per patient. A multivariable model was used to assess the independent effect of enhancing patterns on QSM and R2*, controlling for Expanded Disability Status Scale score, age, and disease duration. Age, Expanded Disability Status Scale score, and disease duration were assessed for the presence of collinearity. All analyses were performed in SAS Version 9.4 (SAS Institute, Cary, North Carolina). P < .05 was considered statistically significant.

**RESULTS**

**Lesion Pattern Characteristics**

A total of 205 white matter lesions (115 enhancing and 90 nonenhancing) were examined in the 64 patients. Two enhancing lesions were excluded because they were too close to the cranium and did not appear intact on QSM. The 203 white matter lesions analyzed consisted of 80 (39.4%) new nodular-enhancing lesions, 33 (16.3%) shell-enhancing lesions, 49 (24.1%) new nonenhancing lesions < 1 year old, and 41 (20.2%) nonenhancing lesions 1–3 years old. Of the 80 new nodular-enhancing lesions, 77 (96.2%) appeared isointense on QSM and 3 (3.8%) lesions were hyperintense on QSM. Of the 33 new shell-enhancing lesions, 30 (90.9%) appeared hyperintense on QSM and 3 (9.1%) lesions were isointense on QSM. All nonenhancing lesions (49 [100%] with age < 1 year and 41 [100%] with age 1–3 years old) were hyperintense on QSM. Example images are shown in Figs 2–5.

**Lesion Susceptibility and R2* Value Analyses**

The reference areas on the contralateral site of 5 new lesions (3 nodular-enhancing, 1 shell-enhancing, and 1 nonenhancing) were covered by MS lesions, which prevented us from assessing reference values for susceptibilities and R2*. The relative susceptibilities and R2* values were measured and calculated in 198 lesions (77 nodular-enhancing, 32 shell-enhancing, 48 nonenhancing < 1 year, and 41 nonenhancing with age 1–3 years).

Quantitative susceptibilities and R2* values are plotted in Fig 6 and are summarized in Table 1. Nodular-enhancing lesions had relative susceptibilities near zero (0.5 ± 4.4 parts per billion), shell-enhancing lesions had increased relative susceptibilities (10.2 ± 5.4 parts per billion), nonenhancing lesions < 1 year old had higher relative susceptibilities (20.2 ± 7.8 parts per billion), and nonenhancing lesions 1–3 years old had the highest relative susceptibility (33.2 ± 8.2 parts per billion). In the generalized estimating equation model, after controlling for Expanded Disability Status Scale score, age, and disease duration, there were significant differences between each 2 types of lesions (P < .0001). The nodular-enhancing, shell-enhancing, and new nonenhancing lesion types (< 1 year) had lower relative susceptibilities compared with the more stable lesions (1–3 years). These relationships were significant at all levels (P < .0001). R2* values relative to normal-appearing WM were −5.6 ± 2.9 Hz, −8.0 ± 2.6 Hz, −3.1 ± 2.3 Hz, and −2.0 ± 2.6 Hz of nodular-enhancing, shell-enhancing, nonenhancing < 1 year old, and nonenhancing 1–3 years old lesions, respectively. These relation-
ships were maintained in the generalized estimating equation model after controlling for the relevant clinical characteristics of age, Expanded Disability Status Scale score, and disease duration. Lesion types of nodular-enhancing, shell-enhancing, and new nonenhancing lesions <1 year old were associated with lower R2* values compared with stable lesions (1–3 years), with shell-enhancing lesions having the largest decrease. All associations were significant at the P < 0.01 level. The results of statistical analysis are shown in Tables 2 and 3.

**DISCUSSION**

Our results demonstrate the following QSM and R2* patterns for MS lesions: nodular-enhancing lesions on T1WI + Gd that are isointense on QSM and hypointense on R2*; susceptibility increases and R2* decreases as lesions develop from nodular- to ring-enhancing; and increase in susceptibility and R2* in lesions that become nonenhancing. According to the imaging physics12 and the known pathophysiology of MS white matter lesions,2 these observed QSM and R2* patterns can be interpreted as sequential pathologic changes in myelin and iron content2 that occur during the early stages of lesion development. Our results suggest that GRE data provide valuable information about MS lesion pathology beyond conventional MR imaging sequences.

We used T1WI + Gd to characterize the temporal sequence of MS lesions, which is important for pathologic interpretation of QSM and R2* patterns. A new MS lesion initially presents as nodular-enhancing on T1WI + Gd. This early stage of active lesion development is characterized by breakdown of the BBB,21 manifesting as enhancement throughout the lesion on T1WI + Gd,20 and myelin damage and uptake by macrophages. The myelin breaking down into debris leads to reduced microstructural heterogeneity, manifesting as R2* hypointensity. At this point, iron does not accumulate in microglia and macrophages, making lesions isointense on QSM. Edema that is present in acute lesions as evidenced as enhancement throughout the lesion on T1WI/Gd.20,23,24 Myelin debris in the lesion’s center is also being further degraded within macrophages, some of which are moving out of the lesion into the peripheral circulation2,25,26 (see myelin content difference between the 2 lesions in Fig 1 in Mehta et al27). This further reduces microstructural heterogeneity and manifests as reduction in R2*. Moreover, loss of diamagnetic lipids increases susceptibility as measured on QSM. Therefore, R2* decrease and QSM increase reflect degradation and removal of myelin debris.

As MS lesions mature into chronic lesions with a closed BBB, iron accumulates predominantly in microglia at the lesion rim, which has been suggested to promote proinflammatory polarization in these cells.27,28 In contrast to demyelination that causes QSM increase and R2* decrease, iron accumulation increases both QSM and R2*. Therefore, simultaneous increase in both QSM and R2* when lesions change from enhancing to nonenhancing may be interpreted as iron accumulation in lesions. There may be slow removal of residual lip degradation products at the same time, but that would only marginally increase QSM and decrease R2*.

It should be noted that BBB recovers at different rates,21,29 which may explain the few outliers that we observed. The QSM hyperintensity in 3 nodular-enhancing lesions may be caused by delayed repair of the BBB or accelerated degradation of myelin. Likewise, the 3 shell-enhancing lesions that are QSM isointense may exhibit fast repair of BBB leakage that leaves no time for susceptibility contents (ie, myelin lipids) to change. Compared with nodular-enhancing lesions, shell-enhancing lesions were captured less frequently in this study and may have more complex structures caused by dynamic pathology.30

MS lesion appearances on QSM and R2* captured at one time point have been reported as heterogeneous or diverse.31 This diversity in susceptibility and R2* values may be caused by the capture of lesions in different stages of development. The dynamic susceptibility change in MS lesion, suggested first by phase imaging32 and cross-section QSM,33 has recently been confirmed in a longitudinal QSM study.34 Our study provides further evidence of dynamic changes in lesion susceptibility and R2*. Here, we attempt to explore the relationship between lesion susceptibility and R2* signal with the underlying pathophysiologic changes that occur over time in MS lesions.2 This provides a biophysically meaningful way to understand lesion diversity, which may be translated into improved patient management.

For example, the data in Fig 6 suggest that 1) T2WI lesions with relative susceptibility value near zero may be nodular-enhancing, opening another venue for the current exploration of identifying enhancing lesions without Gd injection33,35 and 2) simultaneous QSM and R2* increases indicate iron accumulation

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**Table 2: Statistical comparison of relative susceptibility from 4 groups of lesions**

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Nodular</th>
<th>Shell</th>
<th>&lt;1 year</th>
<th>1–3 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shell</td>
<td>−9.8621</td>
<td>[−11.7759, −7.9484], P &lt; .0001</td>
<td>−20.3123</td>
<td>[−23.0281, −17.5959], P &lt; .0001</td>
</tr>
<tr>
<td>&lt;1 year</td>
<td>−10.4502</td>
<td>[−13.6101, −7.2903], P &lt; .0001</td>
<td>−22.7666</td>
<td>[−26.2751, −19.2581], P &lt; .0001</td>
</tr>
<tr>
<td>1–3 years</td>
<td>−12.3164</td>
<td>[−16.3345, −8.2983], P &lt; .0001</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Data given as relative parameter estimates (β). (95% confidence limits). P value. Parameter estimates, 95% CI, and SE from multivariable generalized estimating equation, controlling for age, Expanded Disability Status Scale score, and disease duration. β estimates were obtained by setting the reference category to different lesion group.

**Table 3: Statistical comparison of relative R2* from 4 groups of lesions**

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Nodular</th>
<th>Shell</th>
<th>&lt;1 year</th>
<th>1–3 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shell</td>
<td>2.3409</td>
<td>(1.4980, 3.1837), P &lt; .0001</td>
<td>−4.7717</td>
<td>[−5.7643, −3.7792], P &lt; .0001</td>
</tr>
<tr>
<td>&lt;1 year</td>
<td>−4.6191</td>
<td>[−5.6973, −3.5408], P &lt; .0001</td>
<td>−9.0856</td>
<td>[−10.4502, −7.7208], P &lt; .0001</td>
</tr>
<tr>
<td>1–3 years</td>
<td>−5.9793</td>
<td>[−7.0956, −4.863], P &lt; .0001</td>
<td>−1.7443</td>
<td>[−2.6578, −0.8308], P = .0047</td>
</tr>
</tbody>
</table>

*Data given as relative parameter estimates (β). (95% confidence limits). P value. Parameter estimates, 95% CI, and SE from multivariable generalized estimating equation, controlling for age, Expanded Disability Status Scale, and disease duration. β estimates were obtained by setting the reference category to different lesion group.
in MS lesions, offering an in vivo tool to study iron-related inflammation behind a sealed BBB that cannot be demonstrated with conventional imaging.

There were several limitations in our study. First, this study was not based on longitudinal evaluation of MS lesions but on cross-sectional data. Second, the number of shell-enhancing lesions was low compared with the other types of lesions. Thus, our cross-sectional results warrant future longitudinal studies that may address both limitations.

CONCLUSIONS

Early lesions that are nodular-enhancing on T1WI + Gd show decrease in R2* but no change in QSM, reflecting early breakdown of myelin. As lesions evolve toward shell-enhancing, susceptibility increases and R2* further decreases with progressive degradation and removal of myelin debris. Finally, as lesions become chronic, iron accumulation is demonstrated as both QSM and R2* increase. Thus, combining QSM and R2* can help identify changes in myelin and iron contents as white matter MS lesions develop.


REFERENCES

30. Henderson AP, Barnett MH, Parratt JD, et al. Multiple sclerosis:


