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Published Version
doi:10.1097/WNR.0000000000000244

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An expanded composite scale of MRI-defined disease severity in multiple sclerosis: MRDSS2

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The objective of this study was to test a new version of the Magnetic Resonance Disease Severity Scale (MRDSS2), incorporating cerebral gray matter (GM) and spinal cord involvement from 3 T MRI, in modeling the relationship between MRI and physical disability or cognitive status in multiple sclerosis (MS). Fifty-five MS patients and 30 normal controls underwent high-resolution 3 T MRI. The patients had an Expanded Disability Status Scale score of 1.6 ± 1.7 (mean ± SD). The cerebral normalized GM fraction (GMF), the T2 lesion volume (T2LV), and the ratio of T1 hypointense LV to T2LV (T1/T2) were derived from brain images. Upper cervical spinal cord area (UCCA) was obtained from spinal cord images. A within-subject r-score (difference of MS from normal control) for each MRI component was calculated, equally weighted, and summed to form MRDSS2. With regard to the relationship between physical disability and MRDSS2 or its individual components, MRI–Expanded Disability Status Scale correlations were significant for MRDSS2 (r = 0.33, $P = 0.013$) and UCCA (r = −0.33, $P = 0.015$), but not for GMF (r = 0.198), T2LV (r = 0.707), and T1/T2 (r = 0.240). The inclusion of UCCA appeared to drive this MRI–disability relationship in MRDSS2. With regard to cognition, MRDSS2 showed a larger effect size ($P = 0.035$) than its individual components [GMF ($P = 0.081$), T2LV (P = 0.179), T1/T2 (P = 0.043), and UCCA (P = 0.818)]. In comparing cognitively impaired with cognitively preserved patients (defined by the Minimal Assessment of Cognitive Function in MS). Both cerebral lesions (T1/T2) and atrophy (GMF) appeared to drive this relationship. We describe a new version of the MRDSS, which has been expanded to include cerebral GM and spinal cord involvement. MRDSS2 has concurrent validity with clinical status. NeuroReport 25:1156–1161 © 2014 Wolters Kluwer Health | Lippincott Williams & Wilkins.

Keywords: brain, cognition, MRI, multiple sclerosis, physical disability, spinal cord

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Received 20 June 2014 accepted 14 July 2014

Introduction

Conventional MRI-based brain lesion and atrophy measures have contributed to the understanding of multiple sclerosis (MS) pathophysiology. However, these measures show weak correlations with clinical status, as measured by the Expanded Disability Status Scale (EDSS), and have an unreliable strength for predicting clinical change. Composite MRI measurements offer an emerging approach to assess the full range of MS-related structural changes [1–4].

We previously described a composite scale to define the severity of damage in MS, known as the Magnetic Resonance Disease Severity Scale (MRDSS); this original version (MRDSS1) combined three cerebral measures: (i) T2 hyperintense lesion volume (LV), (ii) the ratio of T1 (hypointense) to T2 LVs (T1/T2), and (iii) normalized whole brain volume (a surrogate of whole brain atrophy). This initial version of the MRDSS showed high effect sizes in comparing MS clinical phenotype groups, was associated with clinical severity measures, and was highly sensitive to longitudinal change when monitoring patients for 3 years [3,4].

Notable limitations are that our previous studies evaluating MRDSS used low-resolution 1.5 T MRI scanning platforms and did not consider cerebral gray matter (GM) or spinal cord damage, both of which are now recognized in numerous studies as key contributors to impairment in patients with MS [5–13]. The goals of this study were (i) to further develop and refine the MRDSS using (a) a 3 T MRI platform with a high-resolution scan protocol and (b) advanced MRI measures – cerebral GM atrophy and spinal...
cord atrophy [new MRDSS version 2 (MRDSS2) = GM volume (GMV) + T2LV + T1/T2 + upper cervical spinal cord area (UCCA)] and (ii) to compare the association of MRDSS1, MRDSS2, and individual MRI measures with neuologic and cognitive functions. This is the first study to consider GM and spinal cord damage in an MS composite scale. We have presented these data in the preliminary form at the 2014 meeting of the American Academy of Neurology, Philadelphia.

Methods
Participants
Table 1 summarizes the demographic and clinical characteristics of the participants. The sample included 55 consecutive patients with MS who met the following criteria: (i) age between 18 and 55 years; (ii) MS diagnosis of either relapsing–remitting, secondary progressive, primary progressive, or clinically isolated demyelinating syndrome [14]; (iii) absence of other major medical, neurologic, or neuropsychiatric disorders; (iv) lack of any relapse or corticosteroid use in the 4 weeks before MRI or start of disease-modifying therapy 6 months before MRI (to reduce confounding effects on MRI); and (v) no history of smoking or substance abuse. Forty-three patients (78%) were receiving disease-modifying treatment at the time of MRI. Within 3 months of MRI, each patient underwent examination by an MS specialist neurologist, with the following pulse sequences (18–20):

1. Brain: coronal three-dimensional modified driven-equilibrium Fourier transform (MDEFT): TR = 7.9 ms, TE = 3.14 ms, flip angle = 15°, slice thickness = 1.6 mm, pixel size = 0.938 × 0.938 mm.
2. Brain: axial T2-weighted fast fluid-attenuated inversion-recovery (FLAIR): TR = 9000 ms, TE = 151 ms, TI = 2250 ms, slice thickness = 2 mm (no gap), pixel size = 0.976 × 0.976 mm.
3. Spine: axial fast spin-echo T2-weighted images: TR = 6117 ms, TE = 110 ms, slice thickness = 3 mm (no gap), pixel size = 0.937 × 0.937 mm.

Image analysis
Brain and spinal cord MRI analysis was carried out in the Laboratory for Neuroimaging Research using Jim (v. 5; Xinapse Systems, Northants, UK, http://www.xinapse.com) and statistical parametric mapping (SPM8; Wellcome Department of Cognitive Neurology, London, UK, http://www.fil.ion.ucl.ac.uk/spm) operating in Matlab (version 2009a; The MathWorks Inc., Natick, Massachusetts, USA). MRI analysts were unaware of clinical information. Our techniques are semi-automated, and we have established their operational procedures and high reliability [12,18,20–23].

Compartment-specific brain volume segmentation
Our pipeline, on the basis of our earlier work [21–23], has been detailed recently [12]. Briefly, an expert first performed manual removal of the skull, paranasal sinuses, and soft tissue overlaying the brain to isolate the intracranial volume (ICV = brain parenchymal tissue + subarachnoid space). Images were then aligned with a common template, bias-field corrected, normalized, and segmented into GM, white matter (WM), and cerebrospinal fluid maps in SPM8. Mutually exclusive masks for each tissue were derived from probability maps. WM volume (WMV), GMV, and brain

Table 1 Demographic and clinical data

<table>
<thead>
<tr>
<th></th>
<th>Multiple sclerosis</th>
<th>Normal controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number (n)</td>
<td>55</td>
<td>30</td>
</tr>
<tr>
<td>Age (years) (mean ± SD)</td>
<td>41.1 ± 9.0</td>
<td>43.9 ± 6.3</td>
</tr>
<tr>
<td>Men [n (%)]</td>
<td>17 (31)</td>
<td>9 (30)</td>
</tr>
<tr>
<td>Disease category [n (%)]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clinically isolated syndrome</td>
<td>4 (7)</td>
<td>–</td>
</tr>
<tr>
<td>Relapsing–remitting</td>
<td>46 (84)</td>
<td>–</td>
</tr>
<tr>
<td>Secondary progressive</td>
<td>4 (7)</td>
<td>–</td>
</tr>
<tr>
<td>Primary progressive</td>
<td>1 (2)</td>
<td>–</td>
</tr>
<tr>
<td>Disease duration (years) (mean ± SD)</td>
<td>8.3 ± 7.4</td>
<td>–</td>
</tr>
<tr>
<td>EDSS score (mean ± SD)</td>
<td>1.6 ± 1.7</td>
<td>–</td>
</tr>
<tr>
<td>Receiving disease-modifying therapy [n (%)]</td>
<td>78%</td>
<td>–</td>
</tr>
</tbody>
</table>

EDSS, Expanded Disease Status Scale.

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parenchymal volume (BPV = WMV + GMV) were derived after manual correction of misclassification of MS lesions and underestimation of the deep GM contour. Normalized compartment-specific global volumes were then obtained as follows: WM fraction (WMF = WMV/ICV), GM fraction (GMF = GMV/ICV), and total brain parenchymal fraction [BPF = (WMV + GMV)/ICV].

Lesion segmentation
Brain FLAIR hyperintense and MDEFT hypointense lesions were expert-segmented using a semi-automated edge-finding tool based on local thresholding in Jim5 to obtain whole brain T2 (FLAIR) hyperintense and T1 (MDEFT) hypointense LVs (T2LV, T1LV), as described previously [18,20]. To assess the destructive potential of lesions, the ratio of T1LV to T2LV (T1/T2) was calculated for each participant.

Spinal cord segmentation
A rapid semi-automated segmentation tool in Jim5 was implemented using the highly reliable and validated active surface method [10] to segment the contour of the spinal cord from the T2-images. The total UCCA, from the top of C2 to the base of C5, was derived for each participant, using consistent landmarks [10].

Creation of MRDSS2
To derive MRDSS2, a $d$-score (difference from NCs) for each MRI component was calculated for all MS patients, as follows:

$$d_{GMF} = \frac{(GMF_{MS} - GMF_{meanNC})}{GMF_{SDMS}}.$$  

The $d$-scores were equally weighted and summed for each patient to form a composite of the four variables as follows:

$$d_{MRDSS2} = \left[ \frac{d_{log(T2LV)} - d_{GMF} - d_{UCCA} + d_{log(T1/T2)}}{4} \right].$$

A composite score using the three original components (used in the previous original version) of MRDSS1 was also calculated for comparison with MRDSS2, as follows:

$$d_{MRDSS1} = \left[ \frac{d_{log(T2LV)} - d_{BPF} + d_{log(T1/T2)}}{3} \right].$$

Thus, two versions of the MRDSS were tested. The second version differed from the first version in two ways: (i) substitution of GMF for BPF; (ii) the addition of spinal cord data. Further, because of the restricted range of the current MS sample, we relied on $d$-scores (rather than $z$-scores, which were used in the original version). Table 2 shows the results of all $d$-scores and MRDSS calculations; Table 3 shows the raw MRI data in the MS and NC groups.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>$d$-Scores for MRI components in the multiple sclerosis group (n = 55)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MRI variable(s)</td>
<td>Mean ± SD (range)</td>
</tr>
<tr>
<td>T2LV (ml)</td>
<td>13.6 (11.7) (2.4 - 1.20)</td>
</tr>
<tr>
<td>T1LV (ml)</td>
<td>0.291 (0.267) (0.0001)</td>
</tr>
<tr>
<td>BPF</td>
<td>0.846 (0.017) (0.0001)</td>
</tr>
<tr>
<td>GMF</td>
<td>0.529 (0.021) (0.14)</td>
</tr>
<tr>
<td>UCCA</td>
<td>2288.2 (322.8) (0.78)</td>
</tr>
</tbody>
</table>

Statistical analysis
The MS and NC groups were compared on all MRI measures using Wilcoxon’s rank sum tests. Associations between MRI-derived data and measures of clinical status were assessed using Spearman’s correlation coefficient in the MS group. Cognitively impaired and cognitively preserved MS groups were compared using two-sample t-tests. In addition to the unadjusted comparison, the association between MRI measures and cognitive impairment was investigated, adjusting for depression (Center for Epidemiologic Studies Depression scale scores) using linear regression. A P-value less than 0.05 was considered significant, and a P-value less than 0.10 was considered a trend to significance in this exploratory study. Effect size ($d$) was also calculated for group comparisons [24].

Results
MRI-disability correlation in the MS group
As shown in Table 4, with regard to all available MRI measures of brain and spinal cord involvement, as well as the two MRDSS versions, we tested their relationship with physical disability (EDSS score). With respect to the individual MRI components that compose the two MRDSS versions, only BPF and UCCA showed significant...
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correlations with EDSS score \( (P < 0.05) \). Considering the two MRDSS versions, MRDSS1 (T2LV, T1/T2, BPF) showed a trend toward a significant correlation with EDSS score \( (P < 0.10) \). In contrast, MRDSS2 (T2LV, T1/T2, GMF, UCCA) showed a significant correlation with EDSS score \( (P < 0.05) \), with the lowest \( P \)-value among all MRI measures; the correlation was only weak-to-moderate in strength. Thus, the addition of GMF and UCCA measures to MRDSS2 appears to improve the validity of the scale from the perspective of overall neurologic disability in patients with MS. Specifically, inclusion of UCCA appeared to be driving this MRI–disability relationship in MRDSS2.

MRI–cognition relationships

As shown in Table 5, with regard to the comparison of MRI measures between cognitively impaired and cognitively preserved patients with MS, we considered the individual components on their own, as well as the two composite scales. With respect to the individual MRI components that composed the two MRDSS versions, only T1/T2 and BPF showed significantly increased severity in the cognitively impaired group \( (P < 0.05) \); GMF showed a trend toward significance \( (P < 0.10) \), and the moderate effect size for this difference approached the level seen for T1/T2 and BPF. Considering the two MRDSS versions, both MRDSS1 (T2LV, T1/T2, BPF) and MRDSS2 (T2LV, T1/T2, GMF, UCCA) showed significantly higher severity in the cognitively impaired group (both \( P \)-s < 0.05), with moderate-to-strong effect sizes \( (0.66–0.74) \). Although the effect size was higher for MRDSS1, the \( P \)-values for both were in the significant range and of similar strengths. Thus, the addition of GMF and UCCA measures to MRDSS2 did not limit the validity of the scale from the perspective of cognitive impairment in patients with MS. Moreover, one notes the value of creating a composite score in that the largest effect sizes for the comparison of the two cognition groups were achieved with the two MRDSS versions versus the individual MRI components. Finally, in considering the relevance of each core component’s contribution to MRDSS2, both cerebral lesions (T1/T2) and cerebral atrophy (GMF) parameters appeared to be driving the relationship between MRDSS2 and cognition.

**Table 4** Relationship between MRI and overall neurologic disability in the multiple sclerosis group \( (n = 55) \)

<table>
<thead>
<tr>
<th>MRI variable(s)</th>
<th>Expanded disability status scale</th>
<th>Spearman’s ( r )</th>
<th>( P )-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>T2LV</td>
<td>0.05</td>
<td>0.707</td>
<td></td>
</tr>
<tr>
<td>T1/T2</td>
<td>0.16</td>
<td>0.240</td>
<td></td>
</tr>
<tr>
<td>BPF</td>
<td>-0.29</td>
<td>0.030*</td>
<td></td>
</tr>
<tr>
<td>GMF</td>
<td>-0.18</td>
<td>0.198</td>
<td></td>
</tr>
<tr>
<td>UCCA</td>
<td>-0.33</td>
<td>0.015*</td>
<td></td>
</tr>
<tr>
<td>MRDSS1 (T2LV, T1/T2, BPF)</td>
<td>0.25</td>
<td>0.067</td>
<td></td>
</tr>
<tr>
<td>MRDSS2 (T2LV, T1/T2, GMF, UCCA)</td>
<td>0.33</td>
<td>0.013*</td>
<td></td>
</tr>
</tbody>
</table>

BPF, brain parenchymal fraction; GMF, gray matter fraction; MRDSS1, previous version of the Magnetic Resonance Disease Severity Scale; MRDSS2, expanded (new) version of the Magnetic Resonance Disease Severity Scale; T1LV, total brain T1 (modified driven-equilibrium Fourier transform) hypointense lesion volume; T2LV, total brain T2 (fluid-attenuated inversion-recovery) hyperintense lesion volume; T1/T2, T1LV/T2LV in each participant; UCCA, upper cervical cord area.

*\( P < 0.05 \).

**Table 5** Comparison of MRI and cognition in the multiple sclerosis group

<table>
<thead>
<tr>
<th>MRI variable(s)</th>
<th>Cognitively impaired ( (n = 20) )</th>
<th>Cognitively preserved ( (n = 35) )</th>
<th>Unadjusted ( P )-value</th>
<th>Adjusted ( P )-value</th>
<th>Effect size ( (d) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( d_{T2LV} )</td>
<td>4.37 (1.08)</td>
<td>3.97 (0.93)</td>
<td>0.171</td>
<td>0.179</td>
<td>0.40</td>
</tr>
<tr>
<td>( d_{T1/T2} )</td>
<td>0.57 (1.07)</td>
<td>0.02 (0.90)</td>
<td>0.043*</td>
<td>0.044*</td>
<td>0.60</td>
</tr>
<tr>
<td>( d_{GMF} )</td>
<td>-0.90 (1.17)</td>
<td>-0.25 (0.82)</td>
<td>0.038*</td>
<td>0.029*</td>
<td>-0.64</td>
</tr>
<tr>
<td>( d_{UCCA} )</td>
<td>-0.65 (1.09)</td>
<td>-0.13 (0.91)</td>
<td>0.081</td>
<td>0.092</td>
<td>-0.52</td>
</tr>
<tr>
<td>MRDSS1 (T2LV, T1/T2, BPF)</td>
<td>1.95 (0.89)</td>
<td>1.40 (0.57)</td>
<td>0.019*</td>
<td>0.011*</td>
<td>0.74</td>
</tr>
<tr>
<td>MRDSS2 (T2LV, T1/T2, GMF, UCCA)</td>
<td>1.38 (0.66)</td>
<td>1.02 (0.41)</td>
<td>0.035*</td>
<td>0.022*</td>
<td>0.66</td>
</tr>
</tbody>
</table>

Values are presented as mean (SD).

BPF, brain parenchymal fraction; GMF, gray matter fraction; MRDSS1, previous version (version 1) of the Magnetic Resonance Disease Severity Scale; MRDSS2, expanded new version of the Magnetic Resonance Disease Severity Scale; T1LV, total brain T1 (modified driven-equilibrium Fourier transform) hypointense lesion volume; T2LV, total brain T2 (fluid-attenuated inversion-recovery) hyperintense lesion volume; T1/T2, T1LV/T2LV in each participant; UCCA, upper cervical cord area.

*\( P < 0.05 \); unadjusted \( P \)-values are based on two-sample \( t \)-tests; depression (Center for Epidemiologic Studies Depression scale)-adjusted \( P \)-values are based on linear regression.

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progress early in the disease course in patients with MS [6]. Whole brain atrophy is dominated by GMV rather than WMV loss [6,8,11,21]. Moreover, the level of global GM atrophy closely tracks MS clinical disease stage/phenotype [8] and has a high degree of relevance in predicting both physical disability [21] and cognitive impairment [22]. One proposed benefit of focusing on GM rather than WM atrophy is that the GM is likely less prone than the WM to transient fluctuations in volume (e.g., fluid and cellular shifts) [6]. However, univariate comparisons in the present study indicate that BPF has a higher validity than GMF for all comparisons. This might be related to the improved reliability in measuring BPF versus GMF because of the more common segmentation misclassifications and variability associated with measurement of the latter [5,18].

A major strength of this study is the inclusion of spinal cord involvement in MRDSS2. Previous MS MRI composite scale versions from our group and other groups have not included spinal cord metrics [1–4]. Yet, a growing body of evidence indicates that spinal cord atrophy is common and highly relevant to disability in advanced stages of MS [9,10,12,13]. Further, spinal cord involvement appears to occur somewhat independently from brain involvement [12], supporting the notion that combining brain and spinal cord MRI metrics provides complementary information on overall disease severity. The inclusion of spinal cord volume in MRDSS2 likely improved the validity of the scale, owing to the strength of its univariate relationship with physical disability (EDSS score).

MRDSS2 takes advantage of the availability and refinement of 3 T MRI, which is growing in use for MS routine clinical care and research evaluations. Our previous work has shown the higher sensitivity to brain lesions and increased relevance toward the prediction of cognitive impairment derived from 3 T versus 1.5 T MRI in MS [20]. This advantage, coupled with the ability to derive higher-resolution images with tolerable scan times, drove our decision to switch the scale to a 3 T platform. However, in the present study, we did not directly compare 1.5 T-derived with 3 T-derived MRDSS scores.

Several additional aspects of our study are worthy of comment. Our MS sample was dominated by mildly affected, treated patients with relapsing forms of the disease. Given that only 9% of our patients had progressive forms of MS, further studies are required to assess the role of this scale in advanced forms of the disease. Because our sample had a restricted range of disease severity, there was limited power to detect disease involvement on each MRI parameter. For example, we did not find significant spinal cord or GM atrophy in the patients relative to controls. Our future studies will test whether methods of normalization of the spinal cord volume [7] and assessment of diffuse pathology in the normal-appearing WM [2] and cortical lesions improve the scale. We will also test nonequal weighting of the MRI measures to improve the validity of the scale. In addition, the T1 hypointense lesions in the present study were defined on gradient-echo rather than spin-echo images; the latter are a more established tool to evaluate destructive lesions [3]. Finally, this cross-sectional study provides the opportunity to determine whether the MRDSS2 predicts the rate of longitudinal clinical deterioration or whether it effectively tracks the response to disease-modifying therapy [25].

Acknowledgements
This study was supported in part by a research grant to Dr Bakshi from the National Multiple Sclerosis Society (RG 4354-A-2).

Conflicts of interest
There are no conflicts of interest.

References


