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## Brain Volume and Diffusion Markers as Predictors of Disability and Short-Term Disease Evolution in Multiple Sclerosis

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### ORIGINAL RESEARCH

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# Brain Volume and Diffusion Markers as Predictors of Disability and Short-Term Disease Evolution in Multiple Sclerosis

**BACKGROUND AND PURPOSE:** MRI markers of neuroaxonal damage in MS have emerged as critical long-term predictors of MS-related disability. Here we investigated the potential of whole-brain diffusivity and brain volume for the prediction of cross-sectional disability and short- to medium-term clinical evolution.

**MATERIALS AND METHODS:** In this multimodal prospective longitudinal MRI study of 54 patients with MS (87% under immunomodulatory therapy, baseline and follow-up at a median of 12 months), ADC histogram analysis, WM lesion load, BPF, whole-brain atrophy rate, MSFC score, and EDSS score were obtained. A total of 44 patients with no relapse at both time points were included.

**RESULTS:** At both time points, ADC histogram analysis provided robust predictors of the MSFC scores (maximal  $R^2 = 0.576$ , P < .001), incorporated cognition and fine-motor skill subscores, and EDSS scores. Significant changes beyond physiologic age-related changes at follow-up were noted for ADC histogram markers and BPF. Stronger diffusivity alterations and brain volume at baseline predicted MSFC decline, as demonstrated by multiple linear regression analysis (mean ADC,  $R^2 = 0.203$ ; P = .003) and lower baseline BPF in patients with declined compared with stable MSFC scores (P = .001). Results were independent of intercurrent relapses.

**CONCLUSIONS:** Diffusion histogram analysis provided stable surrogates of disability in MS and proved sensitive for monitoring disease progression during a median of 12 months. Advanced neuroaxonal pathology at baseline was indicative of an increased risk for sustained progression during a median of 12 months, independent of intercurrent relapses.

**ABBREVIATIONS:** BPF = brain parenchyma fraction; CI = confidence interval; 9-HPT = 9-Hole Peg Test; EDSS = Expanded Disability Status Scale; GM = gray matter; MSFC = MS Functional Composite; PASAT = Paced Auditory Serial Addition Test; PBVC = percentage brain volume change; TWT = timed walk test; WMLL<sub>perc</sub> = WM lesion load volume as percentage of total WM volume

A cute inflammation and demyelination, secondary neuroaxonal pathology, and additional neurotrophic disturbances conjointly lead to clinical impairment in MS.<sup>1,2</sup> Among these factors, the cumulative neuroaxonal damage is a particularly strong determinant of disability.<sup>3</sup> Exceeding a certain threshold of neuroaxonal damage might accelerate a patient's transition to secondary-progressive MS or "sustained progression."<sup>3</sup> Therefore, MRI techniques that are sensitive to the cumulative neuroaxonal damage such as volumetry<sup>2,4</sup> and DWI<sup>5</sup> warrant further investigation to improve the clinical management of MS.

Brain-volume loss in MS is a multifactorial process that originates from inflammatory focal axonal damage and de-

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pletion of myelin sheaths, secondary neuroaxonal degeneration, and other immunologically triggered neurotrophic disturbances.<sup>2,4</sup> It can be observed across all MS subtypes,<sup>6</sup> even in early stages,<sup>7</sup> and its relation to physical and cognitive disability is generally recognized.<sup>8-10</sup> In relapsing-remitting MS, between 47% and 81% of brain atrophy was ascribed to the previous cumulative gadolinium enhancement.<sup>11</sup> Other studies suggest that brain atrophy is a consequence of diffuse pathology rather than focal lesions.<sup>4,12</sup> In fact, signs of strong tissue destruction may occur during the course of MS despite low cumulative inflammatory activity.<sup>13</sup> The strong clinical relevance of brain volumetry in MS is supported by correlations between baseline brain volume and disability occurring 8 years later<sup>14</sup> and associations between early brain atrophy rates and clinical deterioration.<sup>15</sup>

DWI detects alterations of microscopic diffusion processes in MS due to a variety of factors, including loss of myelin sheaths, loss of axonal membranes, neuronal apoptosis, and gliosis formation.<sup>5</sup> It is now well-established that diffusivity measurements are sensitive to MS-related pathology in brain areas that appear normal on conventional T2- and T1weighted images.<sup>5,16-18</sup> Notably, diffusivity changes parallel grades of axonal pathology in animal models<sup>19</sup> and in humans,<sup>20,21</sup> which might explain correlations of diffusivity markers with patient disability status.<sup>22-24</sup> Serial application of DWI revealed progressive microstructural GM changes in

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Table 1: Demographic, clinical, and therapeutic characteristics of patients and controls

	Patients	Controls
No.	44	54
Age (mean $\pm$ SD) (yr)	37.4 (9.6)	38.2 (9.1)
Men/women	16/28	19/35
Age of onset (mean $\pm$ SD) (yr)	30.2 (10.1)	N/A
Disease duration at baseline (mean $\pm$ SD) (yr)	6.6 (6.3)	N/A
Disease type (RRMS/SPMS/PPMS) <sup>a</sup>	38/4/2	N/A
Immunomodulatory therapy (No.) (%)		N/A
$0 = No therapy^{b}$	7 (15.9)	
$1 = \beta$ -interferon <sup>c</sup>	21 (47.7)	
2 = Glatirameracetate	11 (25.0)	
3 = Immunoglobulins	4 (9.0)	
4 = 3-month methylprednisolone	1 (2.2)	

Note:—NA indicates not applicable; PPMS, primary-progressive MS; RRMS, relapsingremitting MS; SPMS, secondary-progressive MS. <sup>a</sup> No significant differences between patients with RRMS and SPMS/PPMS were detected

for age, age of onset, and disease duration. <sup>b</sup> One patient changed to interferon treatment during the study period.

<sup>c</sup> Three patients changed from interferon to mitoxantrone therapy during the study period.

untreated relapsing-remitting MS,<sup>25</sup> and a good prediction of the potential for the clinical status after 5 years in primary-progressive MS.<sup>26</sup>

Despite the sensitivity of DWI, however, there are scant serial data on longitudinal DWI measures, particularly in treated relapsing-remitting MS and in combination with sensitive clinical monitoring instruments such as the MSFC score.<sup>27</sup> Furthermore, serial studies have either focused on DWI<sup>25,26,28,29</sup> or brain volume measurements,<sup>6,9,11,14,30-36</sup> with only 1 serial study on primary-progressive MS and secondary-progressive MS using both techniques.<sup>37</sup> In this prospective, longitudinal, and multimodal MRI study on patients with MS under treatment, we investigated the potential of whole-brain diffusivity and brain volume for the prediction of cross-sectional disability and short-to-medium-term clinical evolution.

#### **Materials and Methods**

#### Patients with MS, Clinical Evaluation, and Controls

Patients were consecutively recruited from the outpatient clinic and the neurologic ward of the Max Planck Institute of Psychiatry, Munich. They fulfilled the criteria of definite MS according to McDonald et al,<sup>38</sup> with the major proportion classified as relapsing-remitting MS (47 patients) (secondary-progressive MS, 5 patients; primaryprogressive MS, 2 patients). Patients with relapses at baseline or at follow-up (n = 10) were excluded from clinicoradiologic correlation analysis to avoid confounding influences from transient clinical exacerbation,<sup>39</sup> leaving 44 patients for the final analysis (Table 1). Disease duration was estimated from a detailed clinical history and file review. Thirty-seven of 44 patients (84%) received immunomodulatory therapy at study entry (Table 1). At baseline and follow-up after a median of 12 months (median, 371 days; range 308-702 days), the EDSS<sup>40</sup> and MSFC scores,<sup>27</sup> comprising the TWT, 9-HPT, and a 3-second version of the PASAT, were obtained. Patients with  $\geq 1$  relapse during the observation interval were identified for post hoc analyses as specified below. The number of patients with MSFC scores available at both time points varied between 38 and 40 due to (disease-related) dropouts in subtests.

Follow-up MSFC scores were interpolated to a 12-month interval (annualized scores). Clinical progression was parameterized as the difference between baseline and annualized follow-up scores. For the MSFC sum score and subscores, patients with a negative annual change value were assigned to the respective progression group in a first step. Second, to reduce false classification into the MSFC-progression group, we classified 20% of the patients with the lowest progression rates as stable. For the EDSS, an increase of  $\geq$ 0.5 point between baseline and follow-up with confirmation 3 months later was classified as EDSS progression.

For proof-of-concept comparisons and estimation of age effect, an age and sex-matched control group free of neurologic or psychiatric disease underwent the same MRI protocol once (n = 54, Table 1).

The study followed the principles of the Declaration of Helsinki and was approved by the local ethics committee. All participants gave their written informed consent.

#### MR Imaging Acquisition and Postprocessing: Overview

Images were acquired on a clinical 1.5T scanner (Signa Excite; GE Healthcare, Milwaukee, Wisconsin). Sequence details and postprocessing steps are described in the on-line supplemental material. In brief, we extracted the following MRI markers: 1) Whole-brain ADC histograms were calculated; mean, variance, skew, and peak-height values were extracted.<sup>18,41</sup> 2) The BPF (brain parenchyma volume divided by total intracranial volume<sup>42</sup>) was calculated at baseline and follow-up from T2-weighted images with high in-plane resolution and CSF/parenchyma contrast by using. 3) The brain-volume change between baseline and follow-up (PBVC) was calculated by using the SIENA algorithm of the FSL software (http://www.fmrib.ox.ac.uk, version 3.2).43 4) For WM lesion load quantification, multispectral image segmentation based on an expectation maximization algorithm was used.44 5) Axial and coronal postgadolinium images of both time points were screened by 2 raters (F.W., P.G.S.) blinded to patient identity and time points.

#### Statistical Analysis

Baseline MRI measures of all patients were compared with those of controls by using univariate multivariate analysis of covariance based on Wilks  $\lambda$ , covarying for age. Group  $\times$  age interaction effects were explored, and the term was removed from the model if not significant (P > .05).

Follow-up MRI variables (ADC histogram metrics, BPF, and WMLL<sub>perc</sub>) as well as follow-up clinical scores were interpolated to a 12-month interval. Paired tests were used to compare baseline and annualized MRI and parametric clinical markers (Wilcoxon signed rank test for EDSS; *t* test for other variables). The patients' annual atrophy rates as calculated by the SIENA algorithm were compared against zero by using a 1-sample *t* test. Annual change rates and 95% CIs for MRI variables of the control group were estimated by linear regression analysis.

For cross-sectional clinico-radiologic correlations, the Spearman rank correlation tests (for EDSS) and the Pearson partial correlation tests corrected for age (for MSFC scores) were applied to baseline and follow-up values. For the 28 MSFC- and 7 EDSS-related tests, Bonferroni-adjusted significance thresholds were defined ( $0.05/28 \sim 0.0018$  and  $0.05/7 \sim 0.0071$ , respectively) to adjust for explorative testing. For baseline and follow-up MSFC and EDSS, stepwise linear regression analysis (variable entry at P < .05, variable removal at P > .10) was appended to identify independent predictors among the MRI variables. Reported  $R^2$  values represent the proportion of explained variance, adjusted for the entry of multiple regressors.

#### Table 2: Change of neurologic scores during a median of 12 months

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Score	No.	Baseline	Follow-Up <sup>a</sup>	P Value <sup>b</sup>	Split Threshold <sup>c</sup>	Progression	Nonprogression
EDSS (median) (range)	44	2.0 (0-5.5)	2.0 (0-5.5)	0.950	≥0.5°	9	35
MSFC (mean) (SD)	38	0.550 (0.415)	0.549 (0.475)	0.969	<-0.047	13	25
TWT (mean) (SD) <sup>d</sup>	39	-0.431 (0.082)	-0.412 (0.098)	0.105	>0.013	16	23
9-HPT (mean) (SD) <sup>e</sup>	40	0.780 (0.735)	0.736 (0.837)	0.382	<-0.113	15	25
PASAT (mean) <sup>f</sup>	39	0.460 (0.693)	0.531 (0.800)	0.414	<-0.101	8	31

Annualized values for MSFC, raw values for EDSS (for details see "Materials and Methods")

<sup>2</sup> Two-sided Wilcoxon signed rank test (EDSS) and paired Student *t* test (MSFC, annualized follow-up values), respectively. <sup>2</sup> See "Materials and Methods" section for details on threshold definition.

d TWT e 9-HPT

f PASAT

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		Patie	Healthy Controls			
MRI Marker	Baseline (Mean)	Follow-Up (Mean) <sup>a</sup>	Annual Change (Mean)	P Value	Estimated Annual Change	
ADC histogram markers					Bp	95% CI
Mean (10 <sup>-6</sup> m <sup>2</sup> /s)	$0.961 \pm 0.031$	$0.965 \pm 0.031$	$0.004 \pm 0.013$	.035°	0.0009 <sup>d</sup>	0.0001; 0.0016
Variance (10 <sup>-6</sup> m <sup>2</sup> /s)	$0.088 \pm 0.008$	$0.089 \pm 0.007$	$0.001 \pm 0.004$	.099 <sup>c</sup>	0.0002 <sup>d</sup>	0; 0.0005
Skew	$1.023 \pm 0.159$	$0.990 \pm 0.151$	$-0.033 \pm 0.078$	.008 <sup>c</sup>	$-0.0038^{d}$	-0.0075; -0.0002
Peak height (10 <sup>-3</sup> )	$2.644 \pm 0.266$	$2.599 \pm 0.253$	$-0.045 \pm 0.144$	.043 <sup>c</sup>	-0.0028	-0.0101; 0.0046
Peak position ( $10^{-6} \text{ m}^2/\text{s}$ )	$0.763 \pm 0.021$	$0.765 \pm 0.020$	$0.003 \pm 0.013$	.172 <sup>c</sup>	0.0004	0.0001; 0.0010
BPF	$0.815 \pm 0.043$	$0.811 \pm 0.039$	$-0.004 \pm 0.016$	.087 <sup>c</sup>	$-0.0023^{d}$	-0.0033; 0.0013
PBVC (%)			$-0.464 \pm 1.233$	.017 <sup>e</sup>		N/A
WMLL <sub>perc</sub> (%)	$3.800 \pm 2.117$	3.921 ± 2.346	$0.121 \pm 0.557$	.157 <sup>c</sup>		N/A
Patients with enhancing lesions (%)	18.0	20.4	N/A	1.000 <sup>f</sup>		N/A

Note:-N/A indicates not applicable.

<sup>a</sup> Annualized values are given except for assessment of gadolinium enhancement. <sup>b</sup> Unstandardized coefficient of linear regression.

Two-sided paired t test.

<sup>f</sup> Fisher exact test

inificant linear correlation with age in the control group ( $P \le .05$ )

PBVC as estimated by the SIENA method. One-sample t test for comparison against zero.

Prediction of clinical progression was analyzed in 2 ways: 1) Baseline MRI markers were compared between patients with and without MSFC progression by using analysis of covariance (2-level group factor, age as a covariate). The analysis was repeated for patients stratified according to EDSS progression. 2) Stepwise linear regression, by using the annual change of MSFC sum score and subscores as dependent variables and baseline MRI variables (ADC histogram markers, BPF, WMLL<sub>perc</sub>) as predictor variables, was applied. To exclude any influence of relapses during the observation interval, we performed the following post hoc analyses: 1) The proportion of patients with intercurrent relapses was compared between the MSFC progression and nonprogression groups (Fisher exact test). 2) MSFC prediction analyses were repeated after exclusion of patients with intercurrent relapses and for patients with relapsing-remitting MS only.

#### Results

#### Clinical Characteristics and Disease Progression

Clinical and demographic sample information is given in Table 1. Table 2 shows comparisons of clinical baseline and annualized follow-up scores. Subtle yet nonsignificant progression of the MSFC sum score, TWT, and 9-HPT and improvement of the PASAT were noted. Twenty percent of patients showed a confirmed EDSS increase of  $\geq 0.5$  points. The MSFC threshold that stratified patients into MSFCprogression and MSFC-nonprogression groups was -0.047 (13 progressive, 25 nonprogressive patients).

Patient/Control Comparison of ADC Histograms and BPF Significant differences between patients and controls were detected for ADC histogram markers (multivariate analysis of covariance, F = 13.895,  $P < 10^{-6}$ ) with an age covariate effect (F = 4.840, P = .001). Similarly, mean BPF differed significantly between patients (0.815  $\pm$  0.043) and controls (0.837  $\pm$ 0.041) (analysis of covariance, F = 6.592, P = .004) with a covariate effect of age (F = 12.580, P = .001) and a trend group  $\times$  age interaction (*F* = 3.667, *P* = .059) (On-line Table 1). BPF correlated with age in the control group ( $r^2 = 0.269$ ,  $P < 10^{-4}$ ) but not in the patient group ( $r^2 = 0.024, P = .314$ ).

#### Longitudinal Changes of MRI Markers

Changes were most explicit for ADC histogram skew, followed by mean ADC and peak height (Table 3). Mean BPF decreased by 0.42% (P = .087), with a similar yet significant mean change calculated by SIENA (PBVC, -0.46%; P = .017), confirming that SIENA is more sensitive to change compared with subtraction of 2 BPF measurements.43 WMLLperc did not change significantly. Annual change rates of all markers in the patient group were outside the 95% CI of the estimated coefficients of controls and exceeded control values by factors 1.7 (BPF) to 8.7 (ADC histogram skew). Gadolinium enhancement was found in 8 patients (18.2%; median gadoliniumenhancement score, 1.0 [range, 1-2]) at baseline and in 9 patients (20.4%; median gadolinium-enhancement score, 2.0 [range, 1–10]) at follow-up. At both baseline and follow-up,



**Fig 1.** Relationship between ADC histogram peak height and MSFC at follow-up (n = 46, r = 0.699, age-corrected partial r = 0.713, P < .001). Patients with acute relapses at the time of assessment were excluded due to potential distortion of their clinical scores.

the gadolinium-enhancement score had no effect on ADC histogram markers or BPF (P > .05).

#### Interrelation between MRI Markers and Disability

Interrelations between MRI and disability were highly similar at baseline and follow-up (see On-line Table 2 for details on correlations at follow-up). At follow-up, interrelations with diffusion markers were more distinct (r = 0.759 [ADC histogram skew, P < .001], Fig 1) than interrelations with WMLLperc (maximum r = -0.487 for MSFC sum score) or BPF (maximum r = 0.438 for 9-HPT). EDSS correlated with 4 of 5 ADC histogram markers and with WMLL<sub>perc</sub>, but not with BPF.

At baseline, linear regression analysis identified ADC histogram variance as an independent predictor of EDSS ( $R^2 = 0.137$ , P = .008), MSFC ( $R^2 = 0.186$ , P = .003), and 9-HPT ( $R^2 = 0.204$ , P = .001). At follow-up, ADC histogram variance was again identified as a predictor of EDSS ( $R^2 = 0.118$ , P = .013); ADC histogram skew, as a predictor of MSFC ( $R^2 = 0.533$ , P < .001); and PASAT ( $R^2 = 0.364$ , P < .001) and ADC histogram height, as a predictor of TWT ( $R^2 = 0.090$ , P = .032) and 9-HPT ( $R^2 = 0.371$ , P < .001).

#### Prediction of Disease Progression from Baseline MRI Markers

Among the baseline MRI markers, ADC and BPF allow a differentiation between future MSFC progression versus nonprogression (P = .042 and P = .001, respectively, Table 4). The latter result was robust toward the Bonferroni correction ( $P < .05/7 \sim .007$ ) (Fig 2*A*). The 2 patient subgroups did not differ with regard to age, age at onset, disease duration, baseline MSFC, baseline EDSS, and the proportion of patients with a relapse during the observation interval (24% and 22%, Fisher exact test, P = .709). Baseline BPF also differentiated between patients with and without EDSS progression (P =

Table 4: Baseline MRI and clinical markers of patients with and without MSFC progression

	MSFC Nonprogression (Mean)	MSFC Progression (Mean)	Р
No. of patients	25	13	
ADC histogram markers <sup>a</sup>			
Mean (10 <sup>-6</sup> m <sup>2</sup> /s)	$0.956 \pm 0.021$	$0.976 \pm 0.039$	.042
Variance (10 <sup>-6</sup> m <sup>2</sup> /s)	$0.088 \pm 0.008$	$0.092 \pm 0.008$	n.s.
Skew	$1.035 \pm 0.116$	$0.951 \pm 0.193$	n.s.
Peak height (10 <sup>-3</sup> )	2.662 ± 0.241	$2.539 \pm 0.282$	n.s.
Peak position (10 <sup>-6</sup> m <sup>2</sup> /s)	$0.758 \pm 0.017$	$0.768 \pm 0.018$	n.s.
BPF	$0.828 \pm 0.040$	$0.783 \pm 0.027$	.001 <sup>b</sup>
WMLLperc (%)	3.365 ± 1.673	$4.435 \pm 2.228$	n.s.
Clinical variables <sup>c</sup>			
Age	$36.9 \pm 9.2$	$39.5 \pm 11.0$	n.s.
Disease duration	$6.6 \pm 6.6$	$7.5 \pm 7.0$	n.s.
Baseline MSFC	$0.53\pm0.46$	$0.59\pm0.33$	n.s.

Note:----n.s. indicates not significant.

<sup>a</sup> Analysis of covariance with 2-level group factor and age as covariate for MRI variables. <sup>b</sup> P value lower than Bonferroni-corrected .05/7  $\sim$  .007.

<sup>c</sup> Two-sided t tests for comparison of clinical variables.

.038). No significant covariate effect of age was found in any comparison.

When we modeled the MSFC subscore changes as continuous variables, mean ADC emerged as an independent predictor of the annual change of MSFC (F = 10.40,  $R^2 =$ 0.203, P = .003, Fig 2B), 9-HPT (F = 12.40,  $R^2 = 0.226$ , P =.001), and PASAT (F = 4.36,  $R^2 = 0.081$ , P = .044). For a change of TWT, baseline WMLL<sub>perc</sub> emerged as a predictor (F = 5.81, P = .021,  $R^2 = 0.112$ ).

Post hoc, an effect of intercurrent relapses on these results was excluded as follows: 1) Patients with intercurrent relapses showed a lower (absolute) mean annual MSFC decrease (n =4, -0.19 ± 0.11) than patients without relapses (n = 9, -0.28 ± 0.26), excluding the finding that MSFC decline was driven by intercurrent relapses. 2) After exclusion of patients with intercurrent relapses, results were stable for the linear regression on the annual MSFC change (baseline mean ADC, r = 0.487, P = .007) and for the comparison of baseline BPF between progressive and nonprogressive patients (P = .0009). 3) Both results were also stable after restriction to patients with relapsing-remitting MS only (n = 38).

#### Discussion

Neuroaxonal damage in MS is a strong mediator of clinical impairment<sup>1</sup> and critical for the development of sustained progression.<sup>3</sup> This clinico-radiologic study on putative neuroaxonal markers obtained 3 main results: 1) Diffusivity histogram metrics were robust predictors of current disability (MSFC and EDSS). 2) Longitudinal analysis revealed pathologically accelerated changes of diffusivity histogram metrics and whole-brain volume during a median of 12 months, mostly in treated patients with relapsing-remitting MS. 3) Advanced brain atrophy and diffusivity alterations at baseline were associated with MSFC decline, independent of intercurrent relapses. More advanced brain atrophy at baseline was also associated with EDSS decline.

ADC histogram parameters of patients with MS distinctly differed from those of controls, with control values found in the range reported in an earlier normative study.<sup>18</sup> In MS,



**Fig 2.** Association of MSFC change with baseline mean ADC and BPF. *A*, Bar plots depict baseline mean ADC (upper plot) and BPF (lower plot) for the control group (n = 54), patients with stable MSFC (n = 25), and patients with progressive MSFC (n = 13). See "Results" section for statistical details. *B*, Annual MSFC change plotted against baseline mean ADC (r = -0.473, P = .003).

abnormal diffusivity is an established finding detectable in focal lesions (ie, WM areas of T2 hyperintensity), whole WM, whole brain, and normal-appearing WM and GM.<sup>5,18,22,45</sup> Comparisons of ADC values among normal-appearing WM, T1 isointense, and T1 hypointense lesions suggest that ADC might parallel different degrees of axonal pathology.<sup>20,21</sup> Similar interrelations between diffusivity and axonal pathology were reported in animal models of MS.<sup>19</sup> Because axonal injury is considered a key factor of disability in MS,<sup>1,3</sup> diffusion imaging might serve as a source of particularly useful surrogate markers. In this study, ADC histogram metrics indeed correlated with the patient's current disability status both at baseline and follow-up (see On-line Table 2 for cross-sectional correlations). Mean ADC proved robust across most clinical measures, while even higher correlations were obtained by histogram distribution measures (eg, skew), suggesting that distribution measures detect diffuse or multifocal diffusivity changes most sensitively.16,46 As expected, the relationship between the MSFC sum score and WMLL<sub>perc</sub> was weaker, likely because focal T2 hyperintensity is histopathologically unspecific and does not capture diffuse pathology.47 Fine motor skills (9-HPT) and cognition (PASAT) also correlated more strongly with diffusivity measures than with BPF and WMLL<sub>perc</sub>. Interrelations with ambulatory function were generally weaker, most likely due to the lack of a spinal marker.

Brain volume was the second marker on which we focused. Patients showed a reduced baseline BPF compared with matched controls, as reported by Kalkers et al<sup>6</sup> and Rudick et al.<sup>48</sup> The physiologic negative relationship between BPF and age was disrupted in patients because younger patients already showed a low BPF. Most interesting, younger patients (median, 36.4 years) exhibited a shorter disease duration (4.6  $\pm$  4.0 versus 9.4  $\pm$  6.8 years, *P* = .003) compared with the older patients, suggesting that young age at onset could increase the risk of developing brain atrophy. This hypothesis, however, needs support from larger samples. BPF shared only a low proportion of variance with WMLL<sub>perc</sub> (~22%), as reported by Guttman et al<sup>49</sup> and Simon et al,<sup>50</sup> confirming that both the

pathology of the normal WM and cortical processes contribute substantially to brain atrophy.<sup>31,34,51</sup> Moderate correlations between BPF and the MSFC sum scores, as described by Kalkers et al,<sup>52</sup> and with the 9-HPT could be established. No correlation with the PASAT was found, as shown in a previous negative report.<sup>53</sup> So far, significant BPF/PASAT correlations have only been reported for 82 patients with MS of different clinical subtypes<sup>54</sup> and for 45 patients with primaryprogressive MS<sup>10</sup>; however, these patient samples showed a more advanced atrophy.

The longitudinal analysis served to probe whether the proposed markers are suitable to monitor and predict disease progression. In contrast to the clinical ratings, diffusivity measures and BPF showed a significant annual progression that exceeded the values estimated for normal aging. An average annual BPF decrease of approximately 0.4%–0.5% is in line with 0.45% reported for treated relapsing-remitting MS,<sup>33,35</sup> larger than the reported rates in healthy subjects (0.1%– 0.3%),<sup>4</sup> and slightly above 0.36% observed in patients with relapsing-remitting MS with optimally suppressed inflammatory activity.<sup>11</sup>

Eventually, the longitudinal analysis revealed that more advanced brain volume loss and higher mean ADC at baseline were associated with short-term progression, as primarily defined from the MSFC score. When we modeled this progression as a continuous variable, avoiding arbitrary thresholding, baseline mean ADC emerged as an independent predictor. Changes of the 9-HPT and PASAT subscores that generally contribute strongly to the MSFC score<sup>55</sup> were also predictable from baseline mean ADC. The same result pattern emerged when the analysis was restricted to patients with relapsingremitting MS and when patients with intercurrent relapses were excluded. Contrary to BPF and diffusivity measures, WMLL<sub>perc</sub> as a focal disease marker showed no progression and proved a weak predictor of current disability and no predictor of disease progression.

While these findings emphasize the impact of nonfocal diffuse pathology, the specificity of the MRI markers for axonal

pathology cannot be claimed. Postmortem data, for example, have revealed a significant contribution of myelin content to mean diffusivity,<sup>56</sup> and also in our study, mean ADC and BPF shared approximately 34% of variance. Diffuse demyelination, through its effects on myelin volume and microscopic diffusion, might therefore influence both volume and ADC measurements. For brain-volume loss in MS, axonal pathology has been proposed to play a prior role because WM volume comprises more axonal (46%) than myelin volume (24%).<sup>4</sup> In turn, higher sensitivity of diffusivity toward demyelination is suggested by larger covariation between WMLL<sub>perc</sub> as an indicator of focal demyelination and ADC markers (average  $R^2 \sim 32\%$ ) compared with BPF ( $R^2 = 21\%$ ). Statistically, results pointed to a higher sensitivity of whole-brain mean ADC compared with BPF in linear prediction models. This effect was robust toward covarying for BPF and remained significant after recalculation of the histograms at a stricter CSF threshold of  $1.5 \times 10^{-3}$  mm<sup>2</sup>/s (data not shown). The correlation between baseline BPF and disease progression was more robust in that changes of EDSS and MSFC could be predicted, yet it was more nonlinear. Most interesting, such a nonlinear relationship between axonal pathology and secondary progression was hypothesized earlier.<sup>3</sup>

Taken together, we propose that longitudinal results add new evidence for the hypothesis that accumulation of diffuse axonal pathology may (gradually) increase the risk for clinical progression<sup>3</sup> whereby the contribution of other nonfocal processes, including diffuse demyelination, remains to be clarified. With regard to brain-volume loss, reaching the stage of secondary progression is particularly critical because slowing brain atrophy rates by immunomodulatory therapy then becomes difficult.<sup>57</sup> Results also demonstrate that subtle clinical progression can be detected by the MSFC score before conventional criteria of secondary disease progression<sup>58</sup> apply. No predictive value of central atrophy for EDSS progression during 14 months was found in a larger study<sup>36</sup>; however, central atrophy is different from BPF used in this study.

Several limitations of the present study need to be considered. Foremost, the small sample size and the observational design impose limitations on result generalizability. In particular, results are not representative of the spontaneous course of MS because treatment was applied as clinically required. Furthermore, intrinsic to the study design, EDSS or MSFC score progression may reflect a combination of sustained progression as attributable to the natural course of disease and nonresponse to therapy. Technically, higher spatial resolution and fully automated repositioning tools would be preferred, particularly for optimal lesion volumetry. Last, for the generation of ADC histograms, CSF masking was based on a previously reported<sup>41</sup> fixed ADC threshold, which leads to partial volume effects by macroscopic CSF and hampers a definite attribution of ADC effects to microscopic tissue properties. Indeed, covariation between BPF and mean ADC decreased about linearly when the ADC clipping threshold was lowered. So far, however, it has not been systematically defined which threshold best reflects the true biological correlation between the 2 markers. To categorically avoid partial volume effects, fluid-attenuated DWI may be useful.59

#### Conclusions

Whole-brain diffusivity and whole-brain volume measurements provide clinically valid and sensitive integral markers to monitor cerebral disease burden in MS during a clinically short time interval of approximately 1 year. The association of advanced brain volume loss and diffusivity changes at baseline with short-term disease progression further suggests that advanced neuroaxonal damage represents a risk for sustained progression.

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