Influence of cigarette smoking on white matter in patients with clinically isolated syndrome as detected by diffusion tensor imaging

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PurpOse
Cigarette smoking has been associated with increased occurrence of multiple sclerosis (MS), as well as clinical disability and disease progression in MS. We aimed to assess the effects of smoking on the white matter (WM) in patients with clinically isolated syndrome (CIS) using diffusion tensor imaging.

MethodS
Smoker patients with CIS (n=16), smoker healthy controls (n=13), nonsmoker patients with CIS (n=17) and nonsmoker healthy controls (n=14) were included. Thirteen regions-of-interest including nonenhancing T1 hypointense lesion and perilesional WM, and 11 normal-appearing white matter (NAWM) regions were drawn on color-coded fractional anisotropy (FA) maps. Lesion load was determined in terms of number and volume of WM hyperintensities.

Results
A tendency towards greater lesion load was found in smoker patients. T1 hypointense lesions and perilesional WM had reduced FA and increased mean diffusivity to a similar degree in smoker and nonsmoker CIS patients. Compared with healthy smokers, smoker CIS patients had more extensive NAWM changes shown by increased mean diffusivity. There was no relationship between diffusion metrics and clinical disability scores, duration of the disease and degree of smoking exposure.

Conclusion
Smoker patients showed a tendency towards having greater number of WM lesions and displayed significantly more extensive NAWM abnormalities.
sion tensor changes relate to the duration of disease, clinical disability and degree of smoking exposure.

**Methods**

**Participants**

The local institutional review board approved the study. All participants gave written informed consent. A total of 60 right-handed subjects (33 CIS patients, 27 healthy volunteers; female (F)/male (M) ratio, 37/23; age range, 20–57 years; mean age, 31.8±8.0 years) participated in this study. Thirty-three CIS patients, admitted to our institution between 2012 and 2014 were included in the study. All patients fulfilled the McDonald diagnostic criteria for MS but had only one clinical episode involving optic neuritis, cervical and/or thoracic myelitis or brainstem/cerebellar syndrome. None of the patients converted to clinically definite MS during the study. Twenty-seven healthy volunteers were recruited using e-mail advertisement. Same individuals also constituted the participants of another MRI study on brain volume alterations. Exclusion criteria were presence of any systemic or neurologic disease other than CIS, receiving steroid therapy, having an attack within four weeks of the MRI scan, having a lifetime history of dependence syndrome other than nicotine, neurologic illness other than CIS, and contraindications to MRI.

Two neurologists (S.D., A.T.) with four and 15 years of experience, blinded to the MRI findings, evaluated all patients and control subjects, and recorded expanded disability status scale scores, age at onset of smoking, pack-years of cigarettes and duration of the disease.

The subjects were categorized into four groups according to the smoking habit and presence of CIS: smoker patients with clinically isolated syndrome (CIS; n=16; F/M ratio, 8/8; mean age, 34.3±8.9 years), smoker healthy controls (Hₐₐ; n=13; F/M ratio, 6/7; mean age, 33.7±7.4 years), nonsmoker patients with clinically isolated syndrome (CISₙₙ; n=17; F/M ratio, 13/4; mean age, 29.7±6.9 years) and nonsmoker healthy controls (Hₐₐ; n=14; F/M ratio, 10/4; mean age, 29.8±7.9 years). Nonsmoker participants had smoked no more than five cigarettes in their lifetime.

**Image acquisition**

Imaging protocol included T1-weighted three-dimensional (3D) high resolution images with 0.9 mm isotropic voxels (MPRAGE) (TR/TE, 2600/3.1 ms; matrix, 224×256; NEX, 1; TA, 4.06; number of slices, 176; slice thickness, 1.00 mm; distance factor, 50%; voxel size, 1 mm; in plane resolution, 1 mm) and 3D double inversion-recovery sampling perfusion with application optimized contrast using different flip angle evolution (DIR-SPACE) (TR/TE, 7500/325 ms; matrix, 192×192; NEX, 1; TA, 6.09; number of slices, 144; slice thickness, 1.12 mm without intersection gap; voxel size, 1 mm; in plane resolutions, 1 mm) imaging on 3.0 T MRI scanner (Magnetom, Tim, Siemens). This imaging sequence was selected because it enables confident lesion detection and delineation by multplanar imaging and cerebrospinal fluid-attenuation (12, 13). Patients received intravenous gadopentetate dimeglumine (0.01 mmol/kg) contrast material. Postcontrast axial T1-weighted 3D MPRAGE coronal fat saturated images (TR/TE, 550/15 ms; matrix, 256×256; NEX, 2) were obtained at 5 min after injection in all patients to detect active demyelinating lesion.

Isotropic high-resolution diffusion tensor imaging (DTI) of the whole brain (single-shot EPI; TR/TE, 8020/83 ms; maximum b value, 1000s/mm²; 60 independent directions; FOV, 256 mm; matrix, 128×128; 64 axial sections with 2 mm thickness without intersection gap; voxel size, 2×2×2 mm) was also selected to recognize the WM integrity.

**Image processing and analysis**

Immediately after image acquisition, a neuroradiologist with 14 years of experience (K.K.O.) reviewed the images to rule out any non-demyelinating lesion or to detect active demyelinating lesions. From this review, the investigator checked the radiologic eligibility of the patients for the study.

To determine lesion load in terms of number and volume, BrainVoyager QX 2.6 for Linux (http://www.brainvoyager.com) was used. Two radiologists with 5 and 14 years of experience (G.D. and K.K.O., respectively) recorded the coordination of every hyperintense WM lesion on 3D DIR-SPACE imaging in consensus. The lesions were ranged by a bounding box in which all the intensities of voxels were fixed to a threshold for a clear visual distinction of lesions and parenchyma (http://support.brainvoyager.com/volume-space/107-volume-rendering/314-users-guide-masking-and-cutting.html). This was done by radiologists in consensus. In this way, the number and total volume of interests were calculated.

FSL (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/) was used for DTI data analysis. All scans were corrected for head motion, and eddy currents using the affine registration. b0 volumes of each subject were extracted and averaged. By fitting a main diffusion tensor in each voxel with FSL DTI Fit tool, the FA and MD maps were calculated. All FA, MD, pre- and postcontrast 3D T1 MPRAGE and DIR-SPACE images were registered to Montreal Neurological Institute template by FLIRT command, a part of FSL. The mean FA and MD of the regions-of-interest (ROIs), specified below were computed for each subject.

A total of 13 ROIs were manually drawn on color-coded FA maps, with a careful review of the co-registered pre- and postcontrast 3D T1-weighted images, 3D DIR-SPACE and T2 turbo spin-echo for accurate localization of ROIs of nonenhancing T1 hypointense lesion without edema in the left frontal WM (if unavailable, then right frontal WM). A perilesional WM 5 mm away from that lesion and 11 NAWM regions of 5–6 mm² were selected. NAWM ROIs were drawn from the genu, splenium and body of the corpus callosum, bilateral corona radiata, superior longitudinal fasciculus, posterior limb of internal capsule, and uncinate fasciculi on color-coded FA maps using Johns Hopkins University WM tractography, and the International Consortium for Brain Mapping DTI-81 WM atlases in FSL. DIR-SPACE sequence was used to avoid lesions in the determination of the NAWM ROIs (Fig. 1).

**Main points**

- Clinically isolated syndrome (CIS), an inflammatory demyelinating disorder of the central nervous system (CNS), is a first clinical episode with features predictive of multiple sclerosis.
- In previous studies, cigarette smoking has been shown to increase the risk of conversion from CIS to MS, worsen clinical disability and exacerbate disease progression.
- Compared with nonsmoker patients, smoker patients with CIS show tendency towards increased white matter lesions, in our study.
- Smoker patients with CIS exhibit more extensive normal-appearing white matter abnormalities than nonsmoker patients on diffusion tensor images.
Statistical analysis

Numerical variables were evaluated for normality of data distribution using the Kolmogorov-Smirnov test. Descriptive statistics were expressed as mean±standard deviation. Independent samples t-test was performed to compare the means of two groups. The Mann Whitney U test was used to compare two groups of nonparametric data. One-way analysis of variance (One-way ANOVA) was performed to compare the differences between groups. Spearman’s rho correlation coefficient was used to examine the relationship between two variables. Multiple linear regression was carried out to assess the association between two or more independent variables and a single continuous dependent variable. The analysis was also used to assess whether confounding exists. The effect of group on dependent variables was examined by multiple linear regression model, after adjustment for age and sex. A P value ≤ 0.05 was accepted as significant. Data analysis was performed by IBM SPSS Statistics 21.0 software (IBM Corp.).

Results

The demographic and clinical features of the patients are summarized in Table 1. There was no significant difference between CIS and HS, nor between CIS nonsmokers and HS nonsmokers in terms of age and sex. Furthermore, age- and sex-corrected analyses were used. Exposure to smoking, which was assessed by mean pack-years, mean cigarettes/day, age at onset of smoking and duration of smoking, was similar in CIS and HS. CIS smoker patients with clinically isolated syndrome; CIS nonsmoker patients with clinically isolated syndrome; HS smoker healthy controls; HNS, nonsmoker healthy controls; EDSS, expanded disability status scale.

Table 1. Baseline demographic characteristics, clinical data, and smoking history

<table>
<thead>
<tr>
<th></th>
<th>CIS S (n=16)</th>
<th>CIS NS (n=17)</th>
<th>H S (n=13)</th>
<th>HNS (n=14)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>34.3±8.9</td>
<td>29.7±6.9</td>
<td>33.7±7.4</td>
<td>29.8±7.9</td>
<td>0.226</td>
</tr>
<tr>
<td>Sex (Female/male), n</td>
<td>8/8</td>
<td>13/4</td>
<td>6/7</td>
<td>10/4</td>
<td>0.223</td>
</tr>
<tr>
<td>Pack-year</td>
<td>9.3±10.1</td>
<td>-</td>
<td>7.7±4.7</td>
<td>-</td>
<td>0.904</td>
</tr>
<tr>
<td>Cigarettes per day</td>
<td>14.3±10.6</td>
<td>-</td>
<td>12.9±6.6</td>
<td>-</td>
<td>0.760</td>
</tr>
<tr>
<td>Duration of smoking (years)</td>
<td>11.6±8.5</td>
<td>-</td>
<td>12.6±8.8</td>
<td>-</td>
<td>0.680</td>
</tr>
<tr>
<td>Age at onset of smoking (years)</td>
<td>20.6±5.4</td>
<td>-</td>
<td>21.0±5.8</td>
<td>-</td>
<td>0.716</td>
</tr>
<tr>
<td>Time lapse from clinical episode (months)</td>
<td>10.8±16.5</td>
<td>9.2±7.0</td>
<td>-</td>
<td>-</td>
<td>0.313</td>
</tr>
<tr>
<td>EDSS</td>
<td>0.06</td>
<td>0.18</td>
<td>-</td>
<td>-</td>
<td>0.600</td>
</tr>
</tbody>
</table>

Data are presented as mean±SD, unless otherwise noted. CISS, smoker patients with clinically isolated syndrome; CISNS, nonsmoker patients with clinically isolated syndrome; HS, smoker healthy controls; HNS, nonsmoker healthy controls; EDSS, expanded disability status scale.
myelinating lesions. ROIs of T1 hypointense lesions and perilesional WM could be placed over the left frontal WM in all patients.

CISₜ had a greater number and volume of WM hyperintense lesions than CISₜₙ on DIR-SPACE imaging (15 vs. 11, \( P = 0.20 \) and 5.2 \( \text{cm}^3 \) vs. 2.4 \( \text{cm}^3 \), \( P = 0.24 \)), although the difference did not reach statistical significance.

Significantly decreased FA and increased MD values were found in perilesional WM and lesions compared with those of NAWM in both CISₜ (mean values: FA=0.33, MD=0.96 for lesions; FA=0.43, MD=0.84 for perilesional WM; FA=0.62, MD=0.77 for NAWM) and CISₜₙ (mean values: FA=0.34, MD=0.98 for lesions; FA=0.47, MD=0.83 for perilesional WM; FA=0.64, MD=0.75 for NAWM) (\( P < 0.05 \), for all). Compared with perilesional WM and NAWM, lesions showed significantly decreased FA and increased MD in CISₜ and CISₜₙ (\( P < 0.05 \), for all). However, FA and MD measures from the lesions and perilesional WM did not differ significantly between CISₜ and CISₜₙ (\( P > 0.05 \), for all) (Fig. 2).

Regions of NAWM with significant FA and/or MD differences in the patient and control groups are presented in Table 2. Compared with Hₜ, CISₜ showed elevated MD in the splenium of corpus callosum, bilateral corona radiate, and superior longitudinal fasciculus (\( P < 0.05 \), for all). Among the nonsmokers, a higher MD in the right corona radiata was found in CISₜₙ than Hₙ (\( P = 0.02 \)). There was no brain region where CISₜ and CISₜₙ had significantly lower MD values than their matched controls. Compared with Hₙₜ, CISₜₙ showed increased FA values in three regions of the NAWM including the body of corpus callosum, left superior longitudinal fasciculus and left posterior limb of the internal capsule (\( P < 0.05 \), for all). None of the NAWM regions showed significant FA change in Hₜ vs. CISₜₙ patients (\( P > 0.05 \), for all).

Compared with Hₙₜ, Hₜ showed no region of FA alteration, while revealing significantly decreased MD values in the left corona radiata and superior longitudinal fasciculus (\( P < 0.05 \), Table 2).

In CIS patients, the mean MD value in the left superior longitudinal fasciculus was significantly higher in smokers than nonsmokers (0.78±0.03×10⁻³ \( \text{mm}^2/\text{s} \) vs. 0.75±0.04×10⁻³ \( \text{mm}^2/\text{s} \), \( P = 0.01 \); Table 2).

Correlation analysis failed to reveal any significant relationship between DTI measures and the duration of disease, expanded disability status scale, and exposure of smoking as assessed by pack-year calculations, cigarettes/day, age at onset, and duration of smoking (\( P > 0.05 \)).

**Discussion**

We aimed to investigate smoking-related WM alterations using DTI in patients with CIS in comparison with healthy controls matched for smoking habit. We calculated FA and MD as the most commonly used DTI measures in chronic T1 hypointense lesions, perilesional WM, and NAWM far from the lesions (14). On ex vivo MRI, chronic, nonenhancing T1 hypointense lesions, so called “black holes,” show hypocellularity, reduced axonal density, decreased myelin and greater matrix disruption compared with non-black hole lesions histopathologically (15). These lesions usually show the greatest WM injury, as also supported by findings of magnetic resonance spectroscopy and magnetization transfer imaging (16). In agreement with previously published reports (5, 17), we found the most significant changes in FA and MD values in T1 hypointense lesions, followed by perilesional WM and NAWM of the patients compared with NAWM of the healthy controls. FA, generally regarded as a measure of WM integrity, is closely related with axonal caliber and the surrounding myelin. MD gives a measure of the average molecular motion depending on cellular size, compartmental fluid imbalance, and degradation products (14, 18). With the abovementioned histopathologic abnormalities in myelin and accompanying primary or secondary axonal loss, reduced FA and elevated MD were expected changes in the WM of our patients with demyelinating disease.

Our study revealed a trend for a greater lesion load on double inversion recovery sequence in smoking patients, although it was not statistically significant. Furthermore, smoker patients had evidently more extensive NAWM changes than nonsmokers (in five ROIs vs. one ROI) compared with matched healthy controls. Additional deteriorating effects of the toxic contents of tobacco smoke such as cyanide, nitric oxide, and free radicals might explain the greater vulnerability of the NAWM in CIS. The associations between thiocyanate, the main metabolite of cyanide and demyelination in the central nervous system of animals (19), and between nitric oxide and axonal degeneration, blocks in axonal conduction, especially in demyelinated axons (20) are well known. Through increased axonal de-
generation, smoking has been blamed for a worse clinical course in MS (21). With no apparent change in FA, decreased MD in the left corona radiata and superior longitudinal fasciculus of Hs may be due to nicotine-related alterations in the WM. The nicotine-induced cytotoxic swelling suggested by Gazdzinski et al. (22) and proinflammatory effects of cigarette smoke on tissue as shown by elevated peripheral leukocytes, and recruitment of polymophonuclear cells, monocytes, macrophages, and increased fibrinogen may be partly responsible for cellular edema and reduced MD levels in the Hs of the current study (23). Some recent studies have hypothesized on the neurotropic and promyelinating effects of smoking and have found higher FA in brain regions such as the prefrontal WM, cingulum, and corpus callosum in smokers compared with nonsmokers (9, 24). Yet, another study on heavy smokers reported decreased FA in the anterior corpus callosum (25). The variable degrees of addiction of smoking in these contradictory research findings. Although limited dependence and exposure to nicotine revealed no significant change, heavy smoking exposure showed marked FA reductions in WM (9, 24). Whether FA or MD is more sensitive and an earlier measure of WM disintegrity remains controversial in the literature (26, 27).

We failed to show a significant correlation of DTI measures with exposure of smoking or duration of the disease in our relatively homogeneous patient group of CIS. This might have resulted from limited cigarette smoking exposure and relatively short duration of disease as well as the younger age of our patients.

Interestingly, Cis in comparison with Hs revealed increased FA in several regions of the NAWM. Increased intracellular water resulting in cellular swelling, fiber reorganization or remyelination can account for an increase in FA (14, 28). Similar findings have been previously shown in the NAWM of patients with chronic recurrent inflammatory optic neuropathy, in T1-hyperintense lesions and normal appearing tissue in patients with MS. The authors attributed these findings to remyelination and glial activation (29, 30). Low expanded disability status scale and absence of conversion to MS during the two years of the study may have been partly related to remyelination in the NAWM, occurring as a compensatory mechanism. More heterogeneous histopathologic processes rather than uniform demyelination or axonal degeneration may be present in the NAWM of patients with CIS.

The limitations of the current study warrant discussion. First, the number of subjects was relatively small. Second, we could not avoid gender disproportion in the smoking and nonsmoking groups because women usually show less tendency to smoke in our population. To minimize this limitation we used age- and sex-corrected analysis. Finally, it was not possible to compare the conversion rate of CIS to MS in smokers vs. nonsmokers due to the relatively short time period of the study. Therefore, studies with more individuals, more detailed characterization of smoking behavior and a longer follow-up are needed.

In conclusion, CIS and smoking appear to cause variable changes on the WM. Smoking patients showed a tendency towards greater numbers of demyelinating WM lesions and significantly more extensive NAWM abnormalities on DTI. Our findings support the adverse effects of smoking on the WM, which in turn may be related with the clinical course of CIS and conversion to MS.

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![Table 2. FA and MD readings in normal-appearing white matter regions across the groups](image)
Yıldray Gökhalk for his help in data acquisition.

**Conflict of interest disclosure**

The authors declared no conflicts of interest.

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