White and Gray Matter Volume Changes and Correlation with Visual Evoked Potential in Patients with Optic Neuritis: A Voxel-Based Morphometry Study

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Background:
The aim of this study was to investigate potential morphological alterations of gray and white matter in patients with optic neuritis (ON) and their relationship with behavioral performance, using voxel-based morphometry (VBM).

Material/Methods:
Twelve (4 males, 8 females) patients with ON and 12 (4 males, 8 females) age-, sex-, and education-matched healthy controls (HCs) underwent magnetic resonance imaging (MRI). Imaging data were analyzed using two-sample t tests to identify group differences in gray and white matter volume (GMV, WMV). Correlation analysis was used to explore relationships between observed GMV and WMV of different areas and visual evoked potential (VEP) in ON.

Results:
Compared with HCs, ON patients had: significantly decreased GMV in the left postcentral gyrus, left inferior frontal gyrus, left anterior cingulate, left and right middle frontal gyrus, and right inferior parietal lobule; decreased WMV in the left middle frontal gyrus, right superior frontal gyrus, left precentral gyrus and right inferior parietal lobule; and increased WMV in the left fusiform gyrus and left inferior parietal lobule. VEP latency of the right eye in ON correlated positively with WMV signal value of the left fusiform gyrus ($r=0.726$, $p=0.008$), and negatively with GMV signal value of the right inferior parietal lobule ($r=-0.611$, $p=0.035$). Duration of ON correlated negatively with WMV signal value of the right superior frontal gyrus ($r=-0.662$, $p=0.019$), while best-corrected visual acuity (VA) of the right eye correlated negatively with WMV signal value of the left middle frontal gyrus ($r=-0.704$, $p=0.011$).

Conclusions:
These results suggest significant brain involvement in ON, which may reflect the underlying pathologic mechanism. Correlational results demonstrate that VEP in ON is closely associated with WMV and GMV atrophy in many brain regions.

MeSH Keywords:
Evoked Potentials, Visual • Functional Neuroimaging • Optic Neuritis • Particulate Matter

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Background

Optic neuritis (ON) refers to inflammation of the optic nerve and is caused by inflammatory demyelination of the optic nerve, by infection, or by non-specific inflammation. The main clinical manifestations include pain during eye movement, sudden vision loss in one or both eyes, visual field defects, a relative afferent pupillary defect, and papilledema [1]. Clinically, ON leads to lesions of the optic nerve axons and apoptosis of retinal ganglion cells. ON can occur in patients with multiple sclerosis (MS) or neuromyelitis optica (NMO). Studies have estimated a prevalence of 5 cases per 100,000 individuals per year in central Europe [2]. Although treatment of the inflammation results in good eyesight recovery in many patients with ON, vision does not return to normal in others, and may be accompanied by abnormal color vision and visual field defects. ON is commonly considered a retinal disease, but a previous study demonstrated that patients with ON presented abnormalities in the visual cortex [3].

Currently, optical coherence tomography (OCT) and visual evoked potential (VEP) are the most important methods for diagnosis of ON. OCT is a noninvasive, high-resolution method that measures the thickness of the retinal nerve fiber layer. A previous study found that patients with MS had thinning of the retinal nerve fiber layer [4]. VEP is an important clinical test and has been used in studying patients with ON and demyelinating diseases [5]. The VEP features of ON are amplitude decreases and prolonged latency, which reflect nerve axon lesions and apoptosis of retinal ganglion cells. VEP can also reflect the severity of optic nerve injury. The significant correlation of multifocal VEP (mfVEP) latency suggests a role for demyelination in promoting axonal loss [6]. In addition, VEP can be used to evaluate the prognosis of ON. A previous study showed that patients with ON still had mfVEP amplitude delays present in many locations, even if they recovered near-normal vision sensitivity [7].

Functional magnetic resonance imaging (fMRI) has been used in ON research. Diffusion tensor imaging can accurately measure fractional anisotropy (FA) and mean diffusivity of the visual pathway. A previous study showed that axial diffusivity (AD) of affected nerves decreases during acute ON and that this AD reduction correlates with the extent of axonal loss [8]. Another study reported that patients with idiopathic demyelinating ON showed decreased mean FA in the affected nerves [9]. In addition, a previous study using fMRI found that patients with ON had reduced functional connectivity within the visual system [10]. Although these findings showed neuronal morphological changes in the ON, there was far less evidence for changes in the neuromechanism of brain in patients with ON.

VBM is a fully automated, whole-brain measurement technique that compares voxel-wise, between-group differences in local brain morphology [11]. The VBM method has been successfully used to investigate mechanisms of diseases such as Alzheimer’s disease [12], obsessive-compulsive disorder [13], and glaucoma [14]. A previous study found that patients with neuromyelitis optica (NMO) showed white matter (WM) atrophy in areas such as the optic chiasm, pons, cerebellum, and frontal gyrus [15]. Another study reported that patients with NMO showed WM atrophy in the corpus callosum and optic radiations, and gray matter (GM) atrophy in the thalamus and prefrontal cortex [16]. Although these findings showed both WM and GM atrophy in patients with ON, few studies have combined the VBM method with VEP. To the best of our knowledge, the current study is the first to explore the changes in WM and GM in ON and their relationship with VEP.

Material and Methods

Subjects

Twelve patients with ON (4 males, 8 females) were recruited from the Ophthalmology Department of the First Affiliated Hospital of Nanchang University. Inclusion criteria were: 1) acute vision loss with or without eye pain; 2) visual field abnormalities associated with damage to nerve fibers; 3) patients with relative pupillary conduction block or abnormal VEP; 4) no clinical or laboratory evidence of compression, ischemic, toxic, genetic, metabolic, or invasive optic neuropathy; 5) no acute vision loss due to retinal disease, alternative eye disease, or disease of the nervous system; 6) no treatment with any drugs before resting-state fMRI scanning; 7) no obvious abnormalities in the brain parenchyma on head MRI scans (including cerebral infarction, cerebral hemorrhage, cerebellar hemangioma, and cerebral tumors); 8) no history of congenital or acquired diseases such as psychiatric disorder, hypertension, diabetes mellitus, or coronary artery disease, and no addictions such as heroin, smoking, or alcohol; 9) no receipt of organ transplantation; and 10) moderate body shape and weight.

Twelve healthy controls (HCs) (4 males, 8 females), who were age-, sex-, and education status-matched to the patients with ON, were also recruited for this study. All HCs met the following criteria: 1) no abnormalities in visual pathways or brain parenchyma on head MRI scans; 2) no ocular disease, naked eye, the corrected visual acuity >1.0; 3) normal nervous system and mental status, with no headaches; and 4) no contraindications for MRI.

The study was authorized by the Ethics Committee of the First Affiliated Hospital of NanChang university. For each subject, the study protocol and procedure were fully explained and consent was obtained. All the methods of this research followed the Declaration of Helsinki and conformed to the principles of medical ethics.
Structural MRI parameters

Participants were scanned on a 3-Tesla MR scanner (Trio, Siemens, Germany) with a 12-channel head coil. High-resolution T1-weighted transversal images covering the whole brain were acquired with a magnetization-prepared rapid gradient echo (MP-RAGE) sequence: 176 slices with section thickness of 1.0 mm; echo time=2.26 ms; repetition time=1900 ms; field of view=215×230 mm. A neuroradiologist evaluated all scans for gross structural abnormalities. No participants were excluded.

Image processing

Functional data were classified by use of MRICro software (www.MRICro.com) to eliminate incomplete data. Structural images were processed with the voxel-based morphometry toolbox (VBM8) (http://dbm.neuro.uni-jena.de/vbm8/) implemented in Statistical Parametric Mapping (SPM8) (Wellcome Department of Imaging Neuroscience, London, UK) running on MATLAB 7.9.0 (R2009b; The Mathworks, Inc, Natick, MA, USA). VBM is a whole-brain, unbiased, semi-automated technique for characterizing regional cerebral differences in structural magnetic resonance images [17]. Segmentation of individual brains into gray matter, white matter, and cerebrospinal fluid was based on VBM8 using the default estimation options (very light bias regularization, 60 mm cut-off for estimating the Gaussian smoothness of bias in image intensity; ICBM [International Consortium for Brain Mapping] European template for initial affine transformation). Spatial normalization into the Montreal Neurological Institute (MNI) standard space was done by the high-dimensional DARTEL (Diffeomorphic Anatomical Registration Through Exponentiated Lie Algebra) approach implemented in VBM8. We used DARTEL to produce gray and white matter templates and used the generated template for all participants’ standardized gray matter and white matter. Finally, the modulated volumes were smoothed with a 6-mm full-width-at-half-maximum (FWHM) Gaussian kernel. Normalized, modulated, smoothed images were submitted to group-level analyses.

Statistical analysis

A general linear model (GLM) analysis was performed with the SPM8 (http://www.fil.ion.ucl.ac.uk/spm) toolkit to investigate the group differences in GM and WM between patients with ON and healthy controls after controlling for the effects of age and sex. The significance level was set at p<0.05, Gaussian random field (GRF) theory corrected, minimum z>2.3. Statistically significant voxels were superimposed on the standardization of 3DT1WI (3-dimensional magnetization prepared rapid acquisition gradient echo sequences) to generate a color drawing. Voxel threshold was selected for 20 neighboring voxels to analyze the area of GM atrophy in patients with ON. For VEP stimulation analysis all subjects underwent pattern-reversal VEP stimulation (RETLPORT electrophysiological instrument, Roland, Brandenburg, Germany) in a dark and quiet room. All participants were in a quiet state. Three active skin electrodes were placed on the scalp along the midline (over the inion) and on lateral positions (right and left). VEP recording was performed at 100-cm distance from the screen. All patients underwent monocular recording with the untested eye covered.

Using stimulus mode with pattern-reversal VEP stimulation, the parameters were set as: stimulus frequency=1.0 Hz and 100 Hz; interphase=500 ms; number of stimulations=100; average screen brightness=5 cd/m²; spatial frequency=50 ms/s; and contrast ratio=90%. Amplitude and latency VEP values were studied at different angular dimensions of the stimulus (120, 60, and 15 degrees for stimuli with small, medium, and large spatial frequencies of stimulation, respectively). VEPs were characterized by a series of N75, P100, and N135 peaks, each characterized by a specific amplitude and latency. The VEP decreased amplitude and prolonged latency, which reflect nerve axon lesions and apoptosis of retinal ganglion cells.

For visual tests analysis we combined the Snellen vision chart with artificial optometry. First, the corneal curvature of all subjects was calculated, then we used the lens measurement instrument to detect the refractive diopter. Second, subjects were kept standing 5 meters from vision chart and the gaze was kept parallel with the vision chart 1.0. Third, according to the diopter number, we chose the best lens to correct the vision. If the subjects had no refractive errors, we recorded the unaided eye vision.

Results

Behavioral results

There were no obvious differences in weight (p=0.749), age (p=0.827), or height (p=0.719) between the patients with ON and the HCs. There were significant differences between the patients with ON and the HCs for best-corrected VA-Right (p<0.001) and best-corrected VA-Left (p=0.001). There were marked differences between the ONs and the HCs in latency of the VEP in the right eye (p=0.001), latency of the VEP in the left eye (p<0.001), amplitudes of the VEP in the right eye (p<0.001), and amplitudes of the VEP in the left eye (p=0.008). Details are presented in Table 1.

Gray and white matter differences

Compared with the HCs, patients with ON had significantly decreased GM volume (GMV) in the brain regions of the left
postcentral gyrus, left inferior frontal gyrus, left anterior cingulate, right middle frontal gyrus, left middle frontal gyrus, and right inferior parietal lobule (Figure 1 [blue], Table 2). Patients with ON also had decreased WM volume (WMV) in the brain regions of the left middle frontal gyrus, right superior frontal gyrus, left precentral gyrus and right inferior parietal lobule, and increased WMV in the left fusiform gyrus and left inferior parietal lobule (Figure 2, Table 3). In addition, we showed the mean of altered GMV values and WMV values between the ON group and HCs (Figure 3).

**Correlation analysis**

In the ON group, the VEP latency of the right eye showed a positive correlation with the WMV signal value of the left fusiform gyrus \((r=0.726, p=0.008)\) and a negative correlation with the GMV signal value of the right inferior parietal lobule \((r=-0.611, p=0.035)\). Duration of ON showed a negative correlation with the WMV signal value of the right superior frontal gyrus \((r=-0.662, p=0.019)\). The best-corrected VA of the right eye showed a negative correlation with the WMVs signal value of the left middle frontal gyrus \((r=-0.704, p=0.011)\). The details are presented in Figure 4.

**Discussion**

To the best of our knowledge, our study is the first to investigate changes in WM and GM in patients with ON using a VBM approach. We found markedly decreased GMV in the brain regions of the left postcentral gyrus, left inferior frontal gyrus, left anterior cingulate, right middle frontal gyrus, left middle frontal gyrus, and right inferior parietal lobule in patients with ON. In addition, we observed that patients with ON had significantly decreased WMV values in the brain regions of the left middle frontal gyrus, right superior frontal gyrus, left precentral gyrus, and right inferior parietal lobule, whereas they had significantly increased WMV values in the cluster of the left fusiform gyrus and left inferior parietal lobule. Furthermore, we observed that the VEP latency of the right eye in patients with ON had a positive correlation with the WMV signal value of the left fusiform gyrus \((r=0.726, p=0.008)\) and a negative correlation with the GMV signal value of the right inferior parietal lobule \((r=-0.611, p=0.035)\). We also found that the duration of ON had a negative correlation with the WMV signal value of the right superior frontal gyrus \((r=-0.662, p=0.019)\). The best-corrected VA of the right eye had a negative correlation with the WMV signal value of the left middle frontal gyrus \((r=-0.704, p=0.011)\). Compared with the HCs, patients with ON had decreased GM volume in several brain regions. The most significant region of atrophy was the left postcentral gyrus. The postcentral gyrus is delimited by the central sulcus anteriorly and the postcentral sulcus posteriorly; this area is the somatosensory cortex in the human brain [18]. Previous studies have identified many diseases that lead to postcentral gyrus dysfunction, such as Parkinson disease [19], Alzheimer disease [20], and multiple sclerosis [21]. Furthermore, Duan et al. [22], using a voxel-based morphometry method, found that patients with NMO had decreased WMV in the right postcentral gyrus. In support of these findings, we also found that patients with ON had significantly decreased GMV in the left postcentral gyrus. We therefore speculate that ON may lead to dysfunction of the left postcentral gyrus.
Figure 1. GMV regional decrease in patients with ON compared with HCs. The significantly decreased regions are seen in the left postcentral gyrus, left inferior frontal gyrus, left anterior cingulate, right middle frontal gyrus, left middle frontal gyrus, and right inferior parietal lobule. (The significance level was set at P<0.05, Gaussian random field (GRF) theory corrected, minimum z >2.3). GMV – grey matter volume; ON – optic neuritis; HCs – health controls.

Table 2. Brain regions with significant differences in grey matter volume between ON group and HCs.

| Condition                  | ON<HC                                                                 |
|----------------------------|----------------------------------------------------------------------|
| Brain areas                | t-score of peak voxel                                               |
|                            | MNI coordinates                                                      |
|                            | Voxel coordinates                                                    |
|                            | X | Y  | Z  |                                             |
| ON<HC                     | 130 | -45 | -15 | 9 | -3.44 |                                        |
| Left inferior frontal gyrus| 105 | -45 | 12  | 27 | -3.95 |                                        |
| Left anterior cingulate    | 52  | -9  | 45  | 15 | -3.35 |                                        |
| Right middle frontal gyrus | 39  | 36  | 18  | 36 | -3.89 |                                        |
| Left middle frontal gyrus  | 91  | -30 | 18  | 54 | -3.48 |                                        |
| Right inferior parietal lobule | 36 | 42  | -48 | 60 | -4.16 |                                        |

Moreover, we found that patients with ON had decreased GMV in the inferior frontal gyrus and bilateral middle frontal gyrus. A previous study demonstrated that the frontal eye fields are causally involved in the attentional top-down control of anticipatory alpha power in the contralateral visual system [23]. In addition, the inferior frontal gyrus is related to emotional and cognitive empathy [24], while the middle frontal gyrus is associated with the processing of language [25]. Many diseases lead to middle frontal gyrus dysfunction, including schizophrenia [26] and attention deficit hyperactivity disorder (ADHD) [27]. Liu et al. [28] found that patients with NMO had significantly increased amplitude of low-frequency fluctuations (ALFF) in the middle frontal gyrus. However, Liang et al. [29] showed that patients with NMO had decreased regional homogeneity (ReHo) in the left medial frontal gyrus. In support of these findings, we found significant regions of atrophy in...
Table 3. Brain regions with significant differences in white matter volume between ON group and HCs.

| Condition                  | ON group and health control | Voxels | MNI coordinates | t-score of peak voxel |
|---------------------------|-----------------------------|--------|-----------------|-----------------------|
| Brain areas               |                             |        |                 |                       |
| ON<HC                     |                             | 31     | –42 57 0        | –3.52                 |
| Left middle frontal gyrus |                             | 35     | 21 60 12        | –4.37                 |
| Right superior frontal gyrus |                         | 50     | –45 0 27        | –3.45                 |
| Left precentral gyrus     |                             | 23     | 39 –42 51       | –3.92                 |
| ON>HC                     |                             | 32     | –51 –57 36      | 3.43                  |

We found that the VEP latency of the right eye in patients with ON correlated negatively with the GMV signal value of the right inferior parietal lobe ($r=-0.611$, $p=0.035$), which agrees with a previous study demonstrating that patients with ON had prolonged latency [30]. The prolonged VEP latency in ON reflects the severity of ON; therefore, the decreased GMV of the inferior frontal gyrus and bilateral middle frontal gyrus in patients with ON. We therefore speculate that ON may lead to dysfunction of the frontal gyrus, while deficit in the frontal gyrus may reflect the damage to eye motion and cognition in patients with ON.
signal values of the right inferior parietal lobule may be related to the severity of ON.

Compared with the HCs, patients with ON had significantly decreased WMV values in the brain regions of the left middle frontal gyrus, right superior frontal gyrus, left precentral gyrus, and right inferior parietal lobule. Many brain function areas are involved in the default mode network (DMN), including the posterior and anterior cingulate cortices, inferior parietal cortex, medial temporal lobes, and medial frontal cortex [31,32]. The DMN is a “resting-state” network, which shows higher activity at rest and tends to have a negative correlation with activity in task-positive networks. Liang et al. [29] found that patients with NMO had decreased ReHo in the left medial frontal gyrus. Duan et al. [22] demonstrated that patients with NMO had decreased WMV in the left middle and medial frontal gyrus, the right superior frontal gyrus, and the bilateral inferior and superior parietal lobules. In support of these findings, we found that patients with ON had decreased WMV values in the left middle frontal gyrus and right inferior parietal lobule, which may reflect dysfunction of the DMN in the patients with ON. Furthermore, the duration of ON correlated negatively with the WMV signal value of the right superior frontal gyrus \((r=-0.662, p=0.019)\), and we speculate that the longer the duration of ON, the more serious the injury to the DMN. We also found that the best-corrected VA of the right eye correlated negatively with WMV signal value of the left middle frontal gyrus \((r=-0.704, p=0.011)\). We speculate that the decreased WMV signal value of the left middle frontal gyrus may reflect the damage to VA in ON.

Interestingly, we found significantly increased WMV values regions in the left fusiform gyrus and left inferior parietal lobule in patients with ON. The fusiform gyrus is the “fusiform face area,” which participates in face processing and social cognition [33,34]. A previous study showed that visual stimuli can cause increases in blood flow in the contralateral posterior fusiform gyrus [35]. Thus, the fusiform gyrus function is closely related to the visual function. The increased WMV areas in the left fusiform gyrus may be related to compensation for the damaged visual function in patients with ON. Furthermore, we observed that the VEP latency of the right eye in ON correlated positively with the WMV signal value of the left fusiform gyrus \((r=0.726, p=0.008)\). The patients with ON had prolongation of VEP latency. The increased VEP latency in ON reflects the severity of ON, and thus the increased WMV values of the fusiform gyrus may relate to the severity of ON.

The inferior parietal lobule contributes to visual word recognition [36]. In our study, we found that patients with ON had higher WMV values in the left inferior parietal lobule and reduced WMV values in the right inferior parietal lobule. Thus, the higher WMV values in the left inferior parietal lobule may reflect functional reorganization to compensate for the damaged areas in patients with ON.

There are some limitations in our study. First, the sample size was too small, which may have affected the accuracy of the results. Second, there were some differences in the duration of ON, and we were not able to entirely separate sex differences and age differences. In future studies we will increase the sample size and choose stricter inclusion criteria.

Figure 3. The mean of altered GMVs values and WMVs values between the ON group and HCs (A, B). GMVs – grey matter volumes; WMVs – white matter volumes; ON – optic neuritis; HCs – health controls.
Figure 4. Correlations between the mean GMV values and WMV values of the different areas and the behavioral performances. The VEP latency of the right eye in ON showed a positive correlation with the WMV signal value of the left fusiform gyrus ($r=0.726, p=0.008$) (A). The VEP latency of the right eye in ON showed a negative correlation with the GMV signal value of the right inferior parietal lobule ($r=-0.611, p=0.035$) (B). The duration of ON showed a negative correlation with the WMV signal value of the right superior frontal gyrus ($r=-0.662, p=0.019$) (C). The best-corrected VA of the right eye showed a negative correlation with WMV signal value of the left middle frontal gyrus ($r=-0.704, p=0.011$) (D). GMV – grey matter volume; WMV – white matter volume; ON – optic neuritis; HCs – healthy controls; VA – visual acuity.

Conclusions

We found that patients with ON had abnormal WM and GM involvement in regional brain changes, which correlated with the VEP latency of the eye. These findings provide important information for understanding the neural mechanisms underlying ON. However, there are some limitations to our study, such as the relatively small sample size, the use of a single center, and the lack of comparison between patients before and after treatment. In future studies we will use other methods to evaluate the changes in brain function in patients with ON.

Disclosure

This was not an industry-supported study. The authors report no conflicts of interest in this work.
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