Abnormal intrinsic functional activity in patients with cervical spondylotic myelopathy: a resting-state fMRI study

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Purpose: We employed resting-state fMRI analyses to reveal central functional reorganization in the brains of patients with cervical spondylotic myelopathy (CSM) and to provide complementary evidence of cortex reorganization in these patients.

Patients and methods: We obtained Fisher’s z transformation amplitude of low-frequency fluctuations (zALFF) and Fisher’s z transformation regional homogeneity (zReHo) measurements from 33 patients with CSM and 33 healthy controls (HC) and used the brain regions with significant alterations in the zALFF or zReHo values as seed regions. Then, we calculated Pearson’s correlation coefficients between the resting-state time courses of each seed and the time series of the rest of the brain. Lastly, we computed correlations between the altered zALFF, zReHo, and functional connectivity with Japanese Orthopaedic Association scores, Neck Disability Index score, and the duration of symptoms in patients with CSM.

Results: zALFF and zReHo values were increased in the left medial superior frontal gyrus (lSFGmed) and left supramarginal gyrus (lSMG) in patients with CSM compared with those in the HC group. Selecting lSFGmed as the seed, we observed increased functional connectivity between it and the left postcentral gyrus (lPoCG) and left rolandic operculum and decreased functional connectivity with the right medial superior frontal gyrus in patients with CSM. In addition, there was a significant increase in the functional connectivity between the lSMG (seed) and the left calcarine and lPoCG in patients with CSM. However, we did not find any significant correlation between the resting-state findings and the clinical performance of patients with CSM.

Conclusion: These observed intrinsic functional changes in the patients with CSM may be related to functional reorganization and reflect the innate cortical plasticity in patients with CSM. Notably, the increased connectivity between the lPoCG and the two seed ROIs indicates the adaptive changes in patients with CSM. These findings provide complementary evidence of cortex reorganization in CSM.

Keywords: ALFF, ReHo, seed-based functional connectivity, cervical spondylotic myelopathy

Introduction

Cervical spondylotic myelopathy (CSM) is the most common disorder of chronic spinal cord compression and is also regarded as a specific incomplete spinal cord injury (SCI). Studies on the local changes in the spinal column or cord of patients with CSM have reported the damage to nerve fibers within the lateral corticospinal tract as the primary cause of the main signs and symptoms observed in patients with CSM. In addition, several studies have reported a degree of functional recovery through cortical...
reorganization and plasticity\textsuperscript{7–9} and central nervous system alterations\textsuperscript{10–14} in patients with CSM. Recently, cerebral functional reorganization or plasticity secondary to neuronal damage in the spinal cord has been accepted as a vital phenomenon in patients with CSM.\textsuperscript{9,10,15} Previous studies\textsuperscript{7,9,16} on sensory-motor cortical plasticity have reported that there is a potential for dynamic reorganization of the brain after secondary injury during the progression of SCI.\textsuperscript{17} There are two major adaptive processes involved in cortical plasticity: 1) synaptic plasticity by modification of preexisting connections and 2) anatomical plasticity through sprouting of axons and dendrites to develop new circuitry.\textsuperscript{16,18} Nowadays, it is considered that the synchronous neural activity of the brain cortices is functionally connected and constitutes a functional network.\textsuperscript{19,20} Given the reported cortical reorganization in patients with complete and incomplete SCI, there is a need to determine whether alterations in intrinsic synchronous functional activity between neurons during resting-state exist, and this may provide information about the complementary functional activity in cortical reorganization. A recent resting-state functional MRI (rs-fMRI) study used the sensorimotor cortex as a priori region of interest (ROI) and found increased amplitude of low-frequency fluctuations (ALFF) in the priori cortex in patients with CSM.\textsuperscript{11} Similarly, another study chose the sensorimotor network as the ROI and observed local neural activity alterations within the priori regions in patients with CSM by measuring regional homogeneity (ReHo) at the resting state.\textsuperscript{12} Moreover, Zhou et al selected the sensorimotor network as the seed region and investigated seed-based functional connectivity strength changes at the voxel level.\textsuperscript{13} They observed altered thalamocortical functional connectivity in two distinct low-frequency bands in patients with CSM.\textsuperscript{14} However, these studies were all hypothesis-driven analyses of some brain region as the specific cortex. Recently, a rs-fMRI study based on data-driven analyses demonstrated that the ALFF/ReHo values in the occipital lobes were decreased and the functional connectivity between the visual cortex and posterior cingulate gyrus was increased in 27 patients with CSM as compared with those of 11 healthy controls (HCs).\textsuperscript{21}

Resting-state fMRI is a powerful and commonly used technique for evaluating spontaneous neural activity.\textsuperscript{22,23} Specifically, ALFF and ReHo are two important data-driven algorithms for the local measurement of spontaneous neural activity. ALFF measures the amplitude of very low-frequency oscillations of the blood oxygen level-dependent (BOLD) signal at the single-voxel level,\textsuperscript{24} while ReHo evaluates the neural synchronization of a given voxel with its adjacent voxels (ie, local neural synchrony).\textsuperscript{25} The combination of the two data-driven analyses without a priori hypothesis may provide a more comprehensive pathophysiological evaluation of brain dysfunction in CSM than each method separately. Moreover, seed-based functional connectivity analysis is a classic approach that correlates the resting-state time courses of a selected ROI (ie, seed) to the time series of the rest of the brain.\textsuperscript{19,20} Exploring the relationship of the synchronous functional activity between adjacent or remote neurons may benefit the understanding of synaptic plasticity and anatomical plasticity in patients with CSM. Therefore, in addition to obtaining ALFF and ReHo values, we conducted seed-based functional connectivity analyses by selecting brain regions with significant ALFF and/or ReHo alterations as the seed regions to obtain more detailed information about the connectivity between these cortices. Subsequently, we assessed the relationship between clinical performance and the altered ALFF, ReHo, and functional connectivity in patients with CSM. Although a previous study\textsuperscript{21} discussed the alterations in resting-state functional activity in patients with CSM, it mainly focused on exploring the relationship of the resting-state-related alternations with visual disorders, and the number of patients with CSM and HCs were not well-matched. Therefore, we aimed to contribute to the current literature on cortical reorganization in patients with CSM by using larger and number-matched study groups in order to identify more elaborate central activity alterations linked to CSM.

\textbf{Materials and methods}

\textbf{Participants}

The present study was approved by the Institutional Review Board of Wuhan University and carried out in accordance with The Code of Ethics of the World Medical Association (Declaration of Helsinki) for experiments involving humans. Thirty-three (17 females; mean age, 54.78±8.41 years [mean ± SD]; range, 42–67 years) patients with CSM were recruited from the Renmin Hospital of Wuhan University through convenience sampling, and 33 (18 females; mean age, 53.52±8.13 years, range, 40–63 years) HCs were enrolled through community health screenings or newspaper advertisements. There were no significant between-group differences in both age and gender. All participants were right-handed and gave written informed consent. Inclusion criterion was definitive CSM, which was based on clear evidence of cord
compression on a cervical spine MRI, such as (1) cervical spondylosis, (2) an ossified posterior longitudinal ligament, or (3) demyelination with hyper-intensity of the cord on T2-weighted images. Meanwhile, two radiologists confirmed spinal cord compression when either the cord surface was clearly indented or the cord diameter had narrowed from compression. Seventeen patients with CSM had right-side lesions, 12 had left-side lesions, and four had bilateral lesions. Patients with more severe signs would be more likely to undergo decompression surgery; therefore, they were excluded from participation. The mean duration of symptoms from disease onset to the date of the MRI examination was 37.0±25.1 months (range, 3 months–8 years). The clinical severity of myelopathy was evaluated using the Japanese Orthopaedic Association (JOA) scores system and Neck Disability Index (NDI) questionnaires. The JOA score system evaluates the severity of myelopathy by assigning scores based on the degree of dysfunction, while the NDI measures the activities of daily living in patients with neck pain. Exclusion criteria included trauma- or infection-related cord compression or other neurological disorders, such as multiple sclerosis, or a history of trauma. Moreover, MRI data from subjects with a head motion >2.0 mm translation or >2.0° rotation in any direction were excluded.

MRI procedure

MRI images were acquired on a 3.0T MR scanner (GE Discovery MR 750) equipped with an eight-channel head coil. A single-shot gradient echo echo-planar imaging) sequence was used to acquire BOLD rs-fMRI data. The scanning parameters were as follows: repetition time (TR) =2 s, echo time (TE) =25 ms, flip angle =90°, slice thickness/gap =3.0/0.6 mm, number of slices =38, field of view (FOV) =24 cm×24 cm, readout bandwidth =250 kHz, and in-plane matrix =64×64. The duration of the resting-state scan was 490 s. During the scan, the subjects were asked to close their eyes and keep their mind blank but avoid falling asleep. For registration purposes, high-resolution anatomical images were acquired from each subject using a sagittal 3D T1-weighted BRAVO sequence with the following parameters: TR =7.2 ms, TE =2.7 ms, inversion time (TI) =450 ms, flip angle =12°, number of slices =160, slice thickness =1.0 mm, FOV =25.6 cm×25.6 cm, readout bandwidth =41.67 kHz, and in-plane matrix =256×256. Sagittal and axial conventional T1-weighted, T2-weighted, and T2-fluid-attenuated inversion recovery (T2-FLAIR) images were acquired from the brain and cervical spinal cord of each patient for diagnosis.

Data preprocessing

All rs-fMRI preprocessing steps were performed using the Data Processing & Analysis of Brain Imaging V2.323 running in MatLab 8.6.0 (Math Works, Natick, MA, USA). Preprocessing involved discarding the first 10 time points to allow the MR signal to reach a steady state and to allow the participants to get used to the scanner environment. Other preprocessing steps included spatial realignment to estimate and correct for subject head motion; slice-timing correction for different acquisition times of slices; and outlier detection, which involved artifact detection tool (ART)-based identification of outlier scans for scrubbing. The high-resolution T1-weighted images were co-registered to the mean realigned fMRI images and segmented into gray matter, white matter, cerebrospinal fluid (CSF), and deformation field images. Next, the fMRI images were spatially normalized to the standard Montreal Neurological Institute (MNI) space and resampled to 3×3×3 mm. Temporal processing involved regressing out confounding factors (including 12 estimated head motion parameters, ART-based scrubbing parameters, and mean signals from white matter and CSF), linear trend removal, and band-pass filtration (0.01–0.08 Hz). Specifically, the mean signals from white matter and CSF were regressed out using the anatomical aCompCor strategy to reduce potentially spurious correlations among distant voxels. Two subjects from each group were excluded due to excessive head motion: one of the two patients with CSM had left spinal cord compression, and the other patient had bilateral spinal cord compression. Thus, 31 subjects from each group were included for further analysis.

ALFF and ReHo calculation

The time series were first smoothed with a Gaussian kernel of 6 mm full-width at half-maximum. The smoothed time series were transformed to frequency domains using the fast Fourier transform to obtain the power spectrum. Subsequently, we calculated the square root at each frequency of the power spectrum and obtained the averaged square root across the 0.01–0.08 Hz frequency range at each voxel. This averaged square root value was chosen as the ALFF value. Finally, this value was transformed using Fisher’s z transformation amplitude of low-frequency fluctuations (zALFF) and used for subsequent group-level analysis.
We obtained ReHo maps in a voxel-wise manner by calculating the Kendall correlation coefficient of a given voxel and its neighboring voxels (26 voxels) from the unsmoothed time series. Each ReHo map was also transformed using Fisher’s z transformation regional homogeneity (zReHo) and used for subsequent group-level analysis.

**Seed-based functional connectivity calculation**

We calculated the Pearson correlation coefficient between the resting-state time courses of the brain regions with significant alterations in the zALFF and/or zReHo (seed regions) and the time series of the rest of the brain. The Pearson correlation coefficient value represented seed-based functional connectivity. Similarly, each Pearson correlation coefficient was transformed using Fisher’s z transformation and used for subsequent group-level analysis.

**Statistical analyses**

**Between-group analyses**

Voxel-wise two-sample t-tests were performed to investigate the differences in the zALFF and zReHo values between patients with CSM and patients in the HC group. Age was included as a nuisance covariate. Type I error was controlled through the use of cluster-wise FDR correction (P<0.01), and the cluster forming threshold was P<0.001. Using the brain regions with significant alterations in the ALFF and ReHo analyses as the seed regions, we performed two-sample t-tests to investigate the between-group differences in each seed-based functional connectivity. Age was also included as a nuisance covariate. We controlled for type I errors through the use of cluster-wise FDR correction (P<0.01), and the cluster forming threshold was P<0.001.

**Correlation analyses**

For each region showing seed-based functional connectivity with a significant between-group difference, we computed the correlation between altered functional connectivity in patients with CSM with the JOA score, NDI score, and duration of symptoms. P<0.01 was set as the statistical significance threshold. In addition, for altered zALFF and zReHo with a significant between-group difference, the same correlation analyses with the three clinical measures (i.e., the JOA score, NDI score, and duration of symptoms) were also conducted.

**Results**

**Demographic and clinical data profiling**

Table 1 and Figure 1 summarize the demographic and clinical data of patients with CSM and patients in the HC group. There were no significant differences in age (P=0.752) or gender (P=0.95) between the two groups. However, there were significant differences in the JOA scores, NDI scores, and duration of symptoms between the two groups (P<0.0001).

### Table 1 Demographic data and clinical measures for the cervical spondylosis myelopathy patients and healthy controls

<table>
<thead>
<tr>
<th>Subject</th>
<th>Cervical spondylosis myelopathy</th>
<th>Healthy controls</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>33</td>
<td>33</td>
<td>n/a</td>
</tr>
<tr>
<td>Age</td>
<td>54.78±8.41</td>
<td>53.52±8.13</td>
<td>0.752</td>
</tr>
<tr>
<td>Gender (male/female)</td>
<td>16/17</td>
<td>15/18</td>
<td>0.95</td>
</tr>
<tr>
<td>Handedness (right/left)</td>
<td>33/0</td>
<td>33/0</td>
<td>n/a</td>
</tr>
<tr>
<td>Laterality of spinal cord compression (right/left/bilateral)</td>
<td>17/12/4</td>
<td>n/a</td>
<td>n/a</td>
</tr>
<tr>
<td>Duration of symptoms (months)</td>
<td>37.0±25.1</td>
<td>n/a</td>
<td>n/a</td>
</tr>
<tr>
<td>JOA scores</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Motor upper</td>
<td>10.53±2.57</td>
<td>17±0</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Motor lower</td>
<td>2.03±0.68</td>
<td>4±0</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Sensory deficit</td>
<td>2.56±1.23</td>
<td>4±0</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Bladder dysfunction</td>
<td>3.13±0.93</td>
<td>6±0</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>NDI scores</td>
<td>2.80±0.20</td>
<td>3±0</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td></td>
<td>0.345±0.102</td>
<td>0.009±0.001</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

**Abbreviations:** JOA, Japanese Orthopaedic Association; NDI, Neck Disability Index.
ALFF and ReHo analyses

Figure 2 shows the differences in zALFF and zReHo values between the two groups. Table 2 lists the Brodmann’s Area (BA) regions, peak MNI coordinates, clusters size, and peak T-value of the differences in the zALFF and zReHo values between the two groups. Compared with patients in the HC group, patients with CSM had significantly increased zALFF in the left medial superior frontal gyrus (ISFGmed) ($P<0.01$, cluster-wise FDR corrected) and significantly increased zReHo in the left supramarginal gyrus (ISMG) ($P<0.01$, cluster-wise FDR corrected) (See Figure 2).

Seed-based functional connectivity analyses

Figures 3 and 4 show the differences in functional connectivity between the two groups when ISFGmed and ISMG were used as the seed ROIs. We observed increased connectivity between the ISFGmed and the left postcentral gyrus (IPoCG) (Figure 3A) and the left rolandic operculum (IROL) (Figure 3B) in patients with CSM as compared to that in the HC group. In addition, connectivity was significantly decreased between the ISFGmed and the right SFGmed (rSFGmed) in the CSM group (Figure 3C). When the ISMG was selected as the seed ROI, we observed increased connectivity between the ISMG and the left calcarine (ICAL) and the IPoCG in patients with CSM (Figure 4D and E). Table 3 lists the BA regions, peak MNI coordinates, clusters size, and peak T-values of the functional connectivity differences between the two groups ($P<0.01$, cluster-wise FDR corrected).
Correlation analyses

Table 4 shows the results of correlation analysis of altered zALFF, zReHo, and functional connectivity with the clinical measures in the CSM group. There were no significant relationships between altered zALFF, zReHo, and functional connectivity with the JOA scores, NDI scores, or duration of symptoms.

Discussion

Through combining both ALFF and ReHo approaches to detect functional plasticity in patients with CSM, as well as investigating whole-brain intrinsic functional connectivity through seed-based functional connectivity analyses, we observed increased zALFF in the lSFGmed and increased zReHo in the lSMG in patients with CSM. Further, we
Figure 4 Functional connectivity differences between the two groups. ISMG selected as seed ROI (P<0.01, cluster-wise FDR corrected). Red-yellow spots, CSM>HC. 
Abbreviations: FDR, false discovery rate; ISMG, left supramarginal gyrus; ICAL, left calcarine; IPoCG, left postcentral gyrus; CSM, cervical spondylotic myelopathy; HC, healthy controls.

Table 3 Functional connectivity differences between the two groups (P<0.01, cluster-wise FDR corrected)

<table>
<thead>
<tr>
<th>Seed ROI</th>
<th>Brain regions</th>
<th>BA</th>
<th>Peak MNI coordinates x, y, z (mm)</th>
<th>Voxels</th>
<th>Peak T value</th>
</tr>
</thead>
<tbody>
<tr>
<td>ISFGmed</td>
<td>IPoCG (CSM &gt; HC)</td>
<td>2/3</td>
<td>-63, -18, 30</td>
<td>76</td>
<td>5.47</td>
</tr>
<tr>
<td></td>
<td>IROL (CSM &gt; HC)</td>
<td>48</td>
<td>-6, -12</td>
<td>41</td>
<td>4.45</td>
</tr>
<tr>
<td></td>
<td>rSFGmed (CSM &lt; HC)</td>
<td>8</td>
<td>6, 27, 48</td>
<td>8</td>
<td>-4.03</td>
</tr>
<tr>
<td>ISMG</td>
<td>ICAL (CSM &gt; HC)</td>
<td>18</td>
<td>-9, -81, 15</td>
<td>113</td>
<td>3.45</td>
</tr>
<tr>
<td></td>
<td>IPoCG (CSM &gt; HC)</td>
<td>2/3</td>
<td>-54, -18, 24</td>
<td>92</td>
<td>4.56</td>
</tr>
</tbody>
</table>

Abbreviations: CSM, cervical spondylotic myelopathy; HC, healthy controls; FDR, false discovery rate; zALFF, Fisher's z transformation amplitude of low-frequency fluctuations; zReHo, Fisher's z transformation regional homogeneity; MNI, Montreal Neurological Institute; ISFGmed, left medial superior frontal gyrus; ISMG, left supramarginal gyrus; IPoCG, left postcentral gyrus.

Table 4 Correlation between altered zALFF, zReHo, and functional connectivity with clinical measures in the CSM group

<table>
<thead>
<tr>
<th>Intrinsic functional changes</th>
<th>JOA scores</th>
<th>NDI scores</th>
<th>Disease duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>zALFF</td>
<td>ISFGmed</td>
<td>-0.198 (0.376)</td>
<td>0.069 (0.752)</td>
</tr>
<tr>
<td>zReHo</td>
<td>ISMG</td>
<td>-0.216 (0.264)</td>
<td>0.143 (0.412)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Seed-based functional connectivity</th>
<th>JOA scores</th>
<th>NDI scores</th>
<th>Disease duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>ISFGmed-IPoCG</td>
<td>-0.163 (0.343)</td>
<td>0.254 (0.271)</td>
<td>0.276 (0.217)</td>
</tr>
<tr>
<td>ISFGmed-IROL</td>
<td>-0.096 (0.564)</td>
<td>0.169 (0.498)</td>
<td>0.047 (0.802)</td>
</tr>
<tr>
<td>ISFGmed- rSFGmed</td>
<td>-0.289 (0.221)</td>
<td>0.037 (0.865)</td>
<td>0.148 (0.399)</td>
</tr>
<tr>
<td>ISMG- ICAL</td>
<td>-0.153 (0.387)</td>
<td>0.294 (0.211)</td>
<td>0.161 (0.495)</td>
</tr>
<tr>
<td>ISMG- IPoCG</td>
<td>-0.173 (0.431)</td>
<td>0.131 (0.523)</td>
<td>0.213 (0.319)</td>
</tr>
</tbody>
</table>

Notes: Values in brackets indicate P-values, and values outside brackets indicate correlation coefficients.
Abbreviations: CSM, cervical spondylotic myelopathy; zALFF, Fisher's z transformation amplitude of low-frequency fluctuations; zReHo, Fisher's z transformation regional homogeneity; JOA, Japanese Orthopaedic Association; NDI, Neck Disability Index; ISFGmed, left medial superior frontal gyrus; rSFGmed, right medial superior frontal gyrus; ISMG, left supramarginal gyrus; ICAL, left calcarine; IROL, left rolandic operculum; IPoCG, left postcentral gyrus.
observed increased functional connectivity between the ISFGmed (seed region) and the lPoCG and the rIOL and decreased functional connectivity between the ISFGmed and the rSFGmed in patients with CSM. In addition, functional connectivity was increased between the ISMG (seed region) and the lCAL and lPoCG in patients with CSM as compared with that in the HC group. These findings on the resting-state-related spontaneous neural activity demonstrate the remarkable brain functional plasticity of patients with CSM. However, we did not find any significant correlations between these altered resting-state-related findings and the clinical performance of patients with CSM.

Increased zALFF and zReHo in patients with CSM
ALFF measures the intensity of spontaneous low-frequency oscillations during the resting state and reveals local cortical intrinsic dynamic functional activity. In the current study, we found that the zALFF was increased in the ISFGmed in patients with CSM. Studies have reported that the SFGmed holds a complete somatotopical representation of body movements through direct connections with the primary motor cortex and spinal cord. As a part of the prefrontal cortex, the SFGmed is also involved in multiple cognitive processes, such as execution of sequential movements, visuomotor association, etc. Recently, many studies have demonstrated cortical reorganization in patients with CSM after spinal cord compression. Specifically, a previous study demonstrated that the areas of cortical representation of the affected limb, such as the adjacent motor territories and the SFGmed cortex, were expanded in patients with CSM. Synaptic plasticity and anatomical plasticity are the two major adaptive processes of cortical reorganization in patients with CSM. The former involves modification of preexisting connections, while the latter involves the development of new circuitry through sprouting of axons and dendrites. The interconnections of axons and dendrites maintain standard cortical representations under the inhibitory influence. However, there may be a disruption of the inhibitory influence in patients with CSM due to afferent or efferent fibers loss. These disinhibited connections between axons and dendrites in patients with CSM facilitate cortical reorganization. ALFF reflects the intrinsic local dynamic activity of neurons, and the observed increase in zALFF in the ISFGmed indicates the increased modulation of cortical activity occurring in patients with CSM. We suspect that this may be attributed to the disinhibitory influence and may be related to cortical plasticity.

ReHo reflects the resting-state neuronal synchronization of intraregional activities. Increased ReHo indicates that the functional activity of a neuron and its adjacent neurons are more temporally synchronous and reflects the perfect coordination of these neurons to achieve a specific function. Recently, studies have shown frequency-dependent changes in the ReHo in different neurologic conditions. According to a previous study, the rs-fMRI signal can be differentiated into four frequency bands: slow 2–5. Unlike the physiological fMRI signal in low frequencies (including slow-5 [0.01–0.027 Hz] and slow-4 [0.027–0.073 Hz]), the contributions of the signal in high frequencies (including slow 2–3 [>0.1 Hz]) to functional connectivity are minor. In this study, we analyzed changes in the ReHo in the frequency range from 0.01 Hz to 0.08 Hz that included the slow-5 and slow-4 but did not separate the two frequency bands. The selected frequency band of the study is in line with the physiological fMRI signal. In the present study, we obtained increased ReHo in the left SMG in patients with CSM. The SMG makes up the anterior part of the inferior parietal lobule (IPL) and consists of a somatosensory-related, higher-order association cortex. Similarly, a recent study that utilized rs-fMRI reported increased ReHo in the right superior parietal lobule in patients with CSM. The SPL and IPL are parietal-integrated regions and are also known as the somatosensory association cortex. Studies have reported that the IPL is related to spatial perception and the interpretation of sensory information and operates as a sensorimotor interface rather than subserving only perceptual functions. Moreover, enhancement activation has been reported in the somatosensory association cortex following the postoperative recovery of function in patients with CSM. A longitudinal fMRI study demonstrated the highly plastic feature of cortical sensorimotor output maps with reorganization in response to changes in the peripheral and central nervous systems. The increased zReHo in the lSMG in patients with CSM may be attributed to functional integration and regulation of injury information from the primary sensory cortex. This may involve greater activity coherence in the sensory-related cortex to compensate for the decreased sensory loss. It has been reported that the clinical signs and symptoms of CSM are not directly consistent with the degree of spinal cord compression and this inconsistency may be explained by the compensatory increase in the ReHo, due to sensory-motor function deficiency in the somatosensory association cortex in patients with CSM.
Further, Bhaqavatula et al. and Duggal et al. have reported that the SFGmed is implicated in cortical reorganization in patients with CSM that involved an increase in the degree of activation in both the precentral gyrus (PrCG) and PoCG. Studies have reported that neuronal plasticity allows neurons to compensate for injury and disease and adjust their activities in response to surrounding changes. For instance, patients who have paralysis may compensate for the impairment of motor function by increased use of muscles unaffected by the disease or trauma, and the functional changes achieved are thought to take place as a result of altered connectivity or neurotransmission within the central nervous system. Previous studies have reported that the SFGmed is implicated in the control of sequential movement and visuomotor association. The PoCG, which belongs to the somatosensory cortex, receives somatosensory inputs from the thalamocortical systems and sends them to other parts of the somatosensory cortex. The increased functional connectivity between the two cortices potentially reflects the compensatory repair of the sensorimotor function in patients with CSM during the recovery stage. An SCI study reported that impaired movement in patients with SCI during the early postinjury stage may result in an overdependence on the associated sensorimotor areas, which play a compensatory role due to the impaired movement function.

In the present study, we also observed increased functional connectivity between the ISFGmed and IROL in patients with CSM. The insula/operculum is involved in interoception and interoceptive awareness and processes signals that are critical for self-awareness. A task-fMRI study observed that the bilateral ROL showed the highest selectivity for bodily self-consciousness based on cardio-visual manipulation and proposed that the ROL processes integrated exteroceptive-interoceptive signals that are necessary for interoceptive awareness and bodily self-consciousness. As mentioned above, the SFGmed holds a complete somatotopical representation of body movements through direct connections with the primary motor cortex and spinal cord. However, a previous magnetic resonance spectroscopy study has reported that there is a decrease in the N-acetylaspartate/creatine metabolite ratio in the primary motor cortex in patients with CSM and cortical levels of N-acetylaspartate/creatine is suggested as a meaningful biomarker in cervical myelopathy, indicative of neuronal damage or dysfunction. Therefore, the SFGmed cortex of patients with CSM may receive less or abnormal information from the primary motor cortex. The enhanced functional connectivity between the ISFGmed and IROL in patients with CSM in the current study may help explain the integration of the exteroceptive-interoceptive signals that are vital to processing bodily self-consciousness due to the deficient information input from the impaired primary motor cortex to resist the body unsteadiness of patients with CSM, which represents an adaptive strategy for functional compensation of patients with CSM.

We found a unique decrease in inter-hemispheric connectivity in the bilateral SFGmed in patients with CSM, which has not been reported before. We suspect that the decreased functional connectivity may be related to the ipsilateral influence on cortex reorganization in patients with CSM, which has been explored in several studies. This influence may allow for high-efficient and low-energy cortical reorganization in patients with CSM. However, further research is required to elucidate what the decreased inter-hemispheric connectivity implies.

The ISMG as the seed ROI

When the ISMG was selected as the seed ROI, we observed increased functional connectivity between the ISMG and IPoCG in patients with CSM. Studies have reported that the ipsilateral PoCG and PrCG seem to be extensively recruited in healthy subjects with increase in the motor task difficulty. Moreover, functional imaging studies have reported enhanced recruitment of the ipsilateral PoCG and PrCG in a wide variety of patients, such as those with stroke, peripheral denervation, and isolated myelitis. The increased functional connectivity between the ISMG and IPoCG is consistent with the suggestion that the brain requires functional rearrangements with fascinating cross-modal plasticity when dealing with sensory loss in patients with CSM. Previous studies on CSM have explored sensory-motor cortical plasticity, which is the dynamic potential of the brain to reorganize following secondary injury in disease progression. Another explanation for the increased functional connectivity between the ISMG and IPoCG in the present study is an analogous cortex
reorganization phenomenon explained in a previous SCI study.16 This previous SCI study reported that compensatory reorganization of somatosensory cortices may play a vital role in limiting the deterioration of motor function and contributing to sensorimotor recovery. Combined with the increased ReHo in the ISMG, the increased association between the ISMG and lPoCG may further compensate for sensorimotor function deficiency, which explains the inconsistency between the degree of spinal cord compression and the clinical signs and symptoms of patients with CSM.

In the current study, we also observed increased functional connectivity between the ISMG and the ICAL. Animal studies have reported that the IPL is involved in the “vestibular cortical circuit” in monkeys.53,54 Meanwhile, studies54–56 have also reported that the SMG is related to vestibular stimulation and involved in the increase in connectivity in patients with bilateral vestibular failure. The SMG contains multisensory neurons that receive visual and/or somatosensory input and also play a role in spatial attention during the control of eye movements.57 The calcarine, which is a sulcus of the medial surface of the occipital lobe, belongs to the primary visual cortex and is associated with visual information processing.58 Increased functional connectivity between the IPL and primary visual cortex has been reported and interpreted as a visual compensatory-dependent mechanism due to the dizziness in vestibular patients.55,56 Studies59,60 on the pathophysiological mechanism of cervical vertigo have indicated that there are many direct neurophysiological connections between the somatosensory, vestibular, and visual systems.61–63 In addition to the common symptoms, including neck pain and stiffness, motor and sensory deficits are also observed. A number of patients with CSM may also experience cervical vertigo. We suggest that the increased functional connectivity between the ISMG and ICAL may result from a similar visual compensatory-dependent mechanism as that seen in vestibular patients. Moreover, a similar resting-state study reported increased functional connectivity between the visual cortex and posterior cingulate lobe in patients with CSM.21 The spinal cord has been reported to have an innate ability to recover varying degrees of sensory, motor, and useful neurological function to accommodate the environmental changes. Thus, the increased association between the ISMG and ICAL, which potentially resulted from the visual compensatory-dependent mechanism, may be in line with the visual sensory recovery in patients with CSM.

**Correlation between altered zALFF, zReHo, and functional connectivity with clinical measures of patients with CSM**

Few fMRI studies on CSM10–13 have reported a significant correlation between functional alterations and JOA scores, NDI scores, or disease duration. Similarly, we did not find any significant correlations between altered zALFF, zReHo, or functional connectivity with these clinical indices. This indicates that the various changes in clinical performance in patients with CSM may be caused by the local insult to the spinal cord rather than intrinsic functional changes of the neurons. Two reasons may explain this phenomenon. First, the interplay between the ongoing destructive mechanisms and the innate reparative processes may eventually reach a balance that implies the innate adaptive or maladaptive plasticity of patients with CSM.12 Next, the JOA and NDI scores are obtained using the questionnaire method; therefore, subjective consciousness and lack of objectivity may influence the scores. Although the JOA system is recommended, it is associated with some clinical disadvantages, including sensitivity, effectiveness, and ignorance of its physical functions for the cervical spine (eg, range of motion of the neck, pain).12,26 Future task-fMRI studies or diffusion tensor imaging studies that could reveal the influence of microstructural white matter changes on cervical cord compression may help identify the elaborate correlation between neuroimage findings and clinical performances of patients with CSM.

Similar to previous fMRI studies, we observed increased zALFF in the ISFGmed and increased zReHo in the IPL in patients with CSM as compared with subjects in the HC group, without concomitant changes in the lPoCG.11–13 It is worth noting that, unlike the previous hypothesis-driven studies,11–13 the present study was based on data-driven methods. A previous study11 that reported increased ALFF in the PoCG selected the sensorimotor cortex (SMC) as an a priori ROI. The researchers calculated the ALFF values within the SMC to obtain individual ALFF maps within a functional SMC mask and compared the group differences between 19 patients with CSM and 19 controls.11 However, we calculated and compared the ALFF difference within the whole brain between 31 patients with CSM and 31 HC. In addition, during the data preprocessing, we regressed out confounding factors, including 12 estimated head motion parameters, ART-based scrubbing parameters, and mean signals from white matter and CSF, which is different from the previous study. Thus, the diverse results may be due to the difference in research approaches, including data preprocessing and subject sizes,
although seed-based functional connectivity analysis revealed a similar increase in the connections between the lPoCG and each of the chosen seed ROIs in the patients with CSM to those that were previously reported.\textsuperscript{7,8,10–13,16,34,40} The observed differences may reflect the adaptive changes in patients with CSM. Interestingly, there is an obvious pattern in our results. Except for the decreased functional connectivity between the rSFGmed and the ISFGmed, all of the other altered functional connectivity occurred in the left hemisphere. Previous studies have explored the ipsilateral characteristics of CSM.\textsuperscript{11–13,16} One of the main interpretations for this is the asymmetrical spinal cord compression or asymmetrical spinothalamic sensory loss.\textsuperscript{13} Other explanations may be the inconsistent maturity of inter-hemispheric regions or the reported ipsilateral recruitment of the PoCG and PrCG for the control of coordinated finger movements during the functional recovery of nonpermanently damaged corticospinal projections in patients with CSM.\textsuperscript{16}

One of the limitations in this study is that we only recruited patients with CSM who had not undergone decompression surgery. Studies have demonstrated that the spinal cord has an innate ability to recover sensory motor function and is helpful in cortical plasticity.\textsuperscript{64} However, decompression surgery in patients with CSM may accelerate the cortical reorganization for recovery of useful neurological functions and optimization of adaptive strategies.\textsuperscript{16,34} Thus, the results of the study may have been improved by enrolling patients before and after decompression surgery. Another limitation is that the rs-fMRI analyses we implemented may have been affected by multiple factors, such as alertness/ sleepiness, substance and medication intake, pain/discomfort, and head motion. Specifically, neck pain is one of the main symptoms of CSM; therefore, the validity of our results may have been affected by uncontrollable head movements caused by discomfort of neck pain during the image acquisition. These confounding factors should be taken into consideration in future studies. Lastly, in our study, Type I error was controlled through the use of cluster-wise FDR correction of $P<0.01$, which is conservative compared to $P<0.001$. We did not observe a significant correlation between the resting-state findings and clinical scores may be potentially associated with the generous threshold. Expanding the sample size and collecting more homogeneous patients with CSM may improve the reliability of the results. In addition, more advanced statistical methods may also contribute to the selection of the most suitable $P$-value.

**Conclusion**

In conclusion, we analyzed the intrinsic neuronal functional activity in the brain and found significant alterations in zALFF, zReHo, and functional connectivity in patients with CSM as compared with those of control participants. These resting-state intrinsic functional changes may be related to functional reorganization and may reflect the innate cortical plasticity in patients with CSM, which indicates the powerful adaptive nature of the brain. Thus, our findings provide complementary evidence regarding cortical reorganization in patients with CSM.

**Abbreviations**

CSM, Cervical spondylotic myelopathy; HC, Healthy Controls; SCI, spinal cord injury; ReHo, regional homogeneity; rs-fMRI, resting-state functional MRI; ALFF, amplitude of low-frequency fluctuations; ISFGmed, left medial superior frontal gyrus; lPoCG, left postcentral gyrus; FDR, false discovery rate; ICAL, left calcarine; lIPL, left inferior parietal lobule; rSFGmed, right medial superior frontal gyrus; PrCG, precentral gyrus; JOA, Japanese Orthopaedic Association; NDI, Neck Disability Index; MNI, Montreal Neurological Institute; BA, Brodmann’s Area; CSF, cerebrospinal fluid; rSPL, right superior parietal lobule; ROI, region of interest; BOLD, blood oxygen level-dependent; FWHM, full-width at half-maximum; T2WI, T2-weighted image; T1WI, T1-weighted image; FOV, field of view; TE, echo time; TR, repetition time; TI, inversion time; T2-FLAIR, T2-fluid-attenuated inversion recovery; ART, artifact detection tool; zALFF, Fisher’s $z$ transformation amplitude of low-frequency fluctuations; zReHo, Fisher’s $z$ transformation regional homogeneity; ROL, Rolandic operculum; DTI, diffusion tensor imaging.

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**Disclosure**

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