Frequency-Specific Alterations of Local Synchronization in Idiopathic Generalized Epilepsy

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Abstract: Recurrently and abnormally hypersynchronous discharge is a striking feature of idiopathic generalized epilepsy (IGE). Resting-state functional magnetic resonance imaging has revealed aberrant spontaneous brain synchronization, predominately in low-frequency range (<0.1 Hz), in individuals with IGE. Little is known, however, about these changes in local synchronization across different frequency bands. We examined alterations to frequency-specific local synchronization in terms of spontaneous blood oxygen level-dependent (BOLD) fluctuations across 5 bands, spanning 0 to 0.25 Hz. Specifically, we compared brain activity in a large cohort of IGE patients (n = 86) to age- and sex-matched normal controls (n = 86). IGE patients showed decreased local synchronization in low frequency (<0.073 Hz), primarily in the default mode network (DMN). IGE patients also exhibited increased local synchronization in high-frequency (>0.073 Hz) in a “conscious perception network,” which is anchored by the pregenual and dorsal anterior cingulate cortex, as well as the bilateral insular cortices, possibly contributing to impaired consciousness. Furthermore, we found frequency-specific alternating local synchronization in the posterior portion of the DMN relative to the anterior part, suggesting an interaction between the disease and frequency bands. Importantly, the aberrant high-frequency local synchronization in the middle cingulate cortex was associated with disease duration, thus linking BOLD frequency changes to disease severity. These findings provide an overview of frequency-specific local synchronization of BOLD fluctuations, and may be helpful in uncovering abnormal synchronous neuronal activity in patients with IGE at specific frequency bands.

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INTRODUCTION

Idiopathic generalized epilepsy (IGE) is a common type of seizure syndrome. Conventionally, generalized seizures are thought to homogeneously affect a large number of brain regions, resulting in abnormal activity across virtually the entire brain. However, focal abnormalities occur in IGE, suggesting local and selectively impaired brain function. Recurrently and abnormally hypersynchronous generalized spike-wave discharges (GSWDs) are a main feature of IGE. Electrophysiological investigations have generally shown that GSWDs are synchronized throughout the brain, resulting in a random organization of the epileptic network. More recently, simultaneous electroencephalography and functional magnetic resonance imaging (EEG-fMRI) studies have demonstrated GSWD-related thalamic activation and default mode network (DMN) deactivation in IGE. In summary, advanced neuroimaging findings regarding (de)activation of certain cortical regions can be interpreted as evidence supporting the focal basis theory of IGE.

Resting-state fMRI (RS-fMRI) constitutes a novel paradigm for investigating spontaneous local brain activity and distant connectivity, and has been widely used to document intrinsic functional abnormalities in various epilepsies. Regional homogeneity (ReHo) analysis, which is a type of RS-fMRI approach, can be used to measure local synchronization in the spontaneous blood oxygenation level-dependent (BOLD) fluctuations of “nearest neighbor” voxels. This local synchronization of BOLD fluctuations is relevant to band-limited power of local field potential. In addition, electrophysiological studies of IGE have shown abnormally hypersynchronous neuronal activity underlying epileptic seizure activity. Accordingly, ReHo analysis-based resting-state fMRI is feasible to detect aberrance of brain temporal synchrony in context of the localization and status of epileptiform activity. Abnormal ReHo is probably associated with changes in neural
activity in certain brain region, and hence may be helpful in localization of the epileptic source in individuals with partial epilepsy.\textsuperscript{15–17} Moreover, we previously reported increased and decreased ReHo in the thalamus and default mode regions, indicating that ReHo might denote abnormal epileptiform activity in generalized tonic-clonic seizures (subsyndrome of IGE) patients.\textsuperscript{15} Note that all the above-mentioned studies predominately detected abnormal epileptic local synchronization in the low-frequency range (<0.1 Hz). Little is known, however, about the changes in local synchronization that occur across different frequency bands in patients with IGE.

Both lower (<0.1 Hz) and higher-frequency (>0.1 Hz) BOLD signals may have specific physiological relevance.\textsuperscript{19,21} For instance, chronic pain studies have specifically demonstrated that aberrant increases in intrinsic brain activity are often derived from higher frequency activity.\textsuperscript{22,23} which corresponds with previous electrophysiological studies reporting the involvement of slow-3, slow-2, and slow-1 frequency bands.\textsuperscript{24,25} Even in the conventional low frequency band (<0.1 Hz), investigator observed stronger BOLD fluctuations at slow-4 (0.027–0.073 Hz) in cortical structures, and at slow-5 (0.01–0.027 Hz) in subcortical structures.\textsuperscript{26} Thus, simply focusing on the conventional low-frequency range runs the risk of overlooking meaningful information associated with frequency-specific spontaneous BOLD fluctuation.\textsuperscript{19,27}

Based on our previous findings regarding frequency-specific amplitude alterations in IGE,\textsuperscript{28} we expect to find disrupted local synchronization of spontaneous BOLD fluctuations across the entire frequency range (within reason) in IGE patients. Thus, we aimed to confirm whether IGE patients exhibit decreased low-frequency synchronization in the DMN based on prior resting-state investigations, and document and spatially localize aberrant high-frequency local synchronization. The purpose of this work was to delineate the whole-brain organization of BOLD local synchronization as a function of a wide range of frequencies and uncover the neurophysiological significance of frequency-specific abnormal synchronous neuronal activity in IGE.

**MATERIALS AND METHODS**

**Participants**

Patients were consecutively recruited at Jinling Hospital, Nanjing, China, between December 2009 and January 2013. Patients were diagnosed according to the criteria of the International League Against Epilepsy (2001). The inclusion criteria for recruitment of patients were as follows: manifestation of typical symptoms of idiopathic generalized tonic–clonic seizures, including tonic extension of the limbs, followed by a clonic phase of rhythmic jerking of the extremities, loss of consciousness during seizures without precursory symptoms of partial epilepsy, and aura; presence of GSWDs in their scalp EEG; no visible focal abnormality in the structural MRI; no history of addiction or neurological disease other than epilepsy; no history of partial seizures; and right handedness. A total of 97 patients fulfilled these inclusion criteria. The exclusion criteria were as follows: self-reported falling asleep during resting-state fMRI scanning, and head motion parameters exceeding 1.5 mm in translation or 1.5° in rotation. After excluding 11 patients owing to excessive head motion, we assessed 86 patients with 3 types of IGE subsyndromes (69 patients with generalized tonic–clonic seizures only [GTCS], 12 patients with juvenile myoclonic epilepsy [JME], and 5 patients with juvenile absence epilepsy [JAE]) in this study. Sixty-four patients took antiepileptic drugs of valproic acid; a part of them additionally took other medications, including topiramate, phenytoin, lamotrigine, and traditional Chinese herb medicine. The demographic and clinical information of these patients is detailed in Table 1.

Eighty-six age- and sex-matched normal controls (NC) were recruited from the staff at the Jinling Hospital. These individuals had no history of neurological disorder or psychiatric illness and no gross abnormalities in brain MRI images. Written informed consent was obtained from all the participants. The study was approved by the Local Medical Ethics Committee at Jinling Hospital, Nanjing University School of Medicine.

**Data Acquisition**

All resting-state fMRI data were acquired using a Siemens Trio 3-Tesla MR scanner located at Jinling Hospital, Nanjing, China. Foam pads and belts were used to minimize head motion. Functional images were acquired using a single-shot, gradient-recalled echo planar imaging sequence (repetition time = 2000 ms, echo time = 30 ms, and flip angle = 90°), which obtained 30 axial slices (field of view = 240 × 240 mm\(^2\), in-plane matrix = 64 × 64, slice thickness/gap = 4 mm/0.4 mm) and 250 functional volumes for each participant, resulting in a total scan time of 500 seconds. Participants were instructed to

| Characteristic                          | IGE (n = 86) | NC (n = 86) | T Value | P Value |
|----------------------------------------|-------------|-------------|---------|---------|
| Sex (male/female)                      | 31/55       | 31/55       |         |         |
| Age, y                                 | 25.77 ± 6.31| 25.01 ± 6.51| 0.7774  | 0.4380  |
| Handedness (right/left)                | 86/0        | 86/0        |         |         |
| IGE subsyndrome (no. of patients)      | GTCS (69), JME (12), JAE (5) | —         |         |         |
| Duration, y                            | 8.93 ± 8.41 | —           |         |         |
| Onset age, y                           | 16.84 ± 7.93| —           |         |         |
| Seizure frequency, times/y             | 18.91 ± 59.19\(^*\) | —         |         |         |

GTCS = generalized tonic–clonic seizures, IGE = idiopathic generalized epilepsy, JAE = juvenile absence epilepsy, JME = juvenile myoclonic epilepsy, NC = normal controls, SD = standard deviation. The P value was obtained from 2-sample t tests.

\(^*\) One patient data missed.
rest with their eyes closed, not think of anything in particular, and not fall asleep.

Data Processing

For functional data processing, we used SPM8 (http://www.fil.ion.ucl.ac.uk/spm) and DPARSF (http://www.restfmri.net). The functional images were corrected for temporal differences by calibrating the slice timing and realigning the images to account for head motion. Functional images were warped into standard stereotaxic space at a resolution of 3 x 3 x 3 mm³ using the Montreal Neurological Institute (MNI) echiplanar imaging template. Next, we used regression processes to remove several sources of spurious variances (6 parameters obtained by rigid-body head motion correction, the time courses averaged over the white matter signal, the cerebrospinal fluid signal, and the global brain signal). Finally, we removed any linear trends. According to the previous electro-physiological22,25 and resting-state fMRI studies,26,28 we used 5 frequency bands: slow-6 (0–0.01 Hz), slow-5 (0.01–0.027 Hz), slow-4 (0.027–0.073 Hz), slow-3 (0.073–0.198 Hz), and slow-2 (0.198–0.25 Hz) to detect frequency-specific local synchronization.12 The center frequencies and intervals of specific frequency bands were based on a previous study.25 Slow-6 occupied the lowest frequency band, which contributed more to ReHo in the putamen.25 Slow-5 and slow-4 mainly represented the frequency range (0.01–0.073 Hz) widely employed in resting-state fMRI studies. Slow-3 represented a combination of the conventional frequency band (0.073–0.1 Hz) and the frequency band discarded through temporal filtering (0.1–0.198 Hz).19 Slow-2 was generally considered to be highly contaminated with respiration signal. However, the BOLD fluctuations in slow-2 frequency band were significantly associated with neuronal fluctuations, and thus may provide more information about functional integration.12 The data from each of 5 frequency-specific ReHo maps for each individual were divided by the global mean ReHo value and then spatially smoothed (full width at half maximum = 6 mm).

Statistical Analysis

We conducted individual 1-sample t tests on the ReHo maps in each group at each frequency band. Thresholds were set at P < 1.00E-07 (uncorrected). To disentangle frequency-specific regional effects, the ReHo maps were entered into a 2-way repeated-measures analysis of variance (ANOVA), with 5 levels (5 bands: 0–0.01; 0.01–0.027; 0.027–0.073; 0.073–0.198; and 0.198–0.25 Hz) for the within-subject factors, and 2 levels (IGE and NC groups) for the between-subject factors. This repeated ANOVA was implemented using the 3dANOVA3 command line in the AFNI toolkit. This model applied an automated anatomical labeling (AAL) mask to each voxel to produce an F-map of the “frequency by group” interaction and an F-map of group main effect. The interaction threshold for the F-maps was set such that the significance level was P < 0.05 (combined false discovery rate [FDR]-corrected height threshold z > 1.96 and extent threshold k > 20 voxels). At first, we used the statistical threshold described above for the main effect analysis. However, no voxel survived at this conservative threshold. Therefore, we used a relatively lenient threshold, P < 0.05 (combined voxel height threshold F > 3.90 and extent threshold k > 141 voxels), with Monte Carlo simulation within the AAL mask for the main effect analysis. Clusters that survived from the group main effect were entered into post hoc 2-sample t tests for each frequency band. The t-maps were set such that the height threshold FDR-corrected significance level was P < 0.05 and the extent threshold was k > 20 voxels.

Head Motion

To moderate the effects of head motion on estimates of ReHo, we first computed the mean framewise displacement (FD) by averaging the FD from every time point for each subject.29 There were significant differences for the mean FD between 2 groups (T = 2.787, P = 0.0059). Second, we used a scrubbing approach to censor “bad” volumes with FD > 0.5 mm to test the robustness of our findings.29

Linear Correlation Between Local Synchronization and Clinical Variables

To investigate the clinical relevance of altered local frequency-specific synchronization in IGE, we correlated the clinical variables (disease duration, onset age, and seizure frequency) with ReHo values using Spearman correlation analysis. Because none of the clinical variables showed a Gaussian distribution, the regions of interest (ROIs) were selected from the clusters showing significantly increased and decreased ReHo values as per the results of post hoc 2-sample t tests. We averaged the ReHo value within each ROI and performed Spearman correlation analysis.

RESULTS

Frequency-Specific Local Synchronization Features

To assess whole-brain full-bandwidth local synchronization profiles of BOLD oscillation during resting-state fMRI, we used 1-sample t tests at each frequency band for each group (Figure 1). In both the groups, visual inspection of the patterns of ReHo in the lower frequency bands (0.01–0.027 and 0.027–0.073 Hz) showed that the DMN regions had higher ReHo values than other brain regions. These reproducible findings were consistent with the previous studies that used typical frequency bands (<0.1 Hz).12 Across the 2 groups, the frequency-specific local synchronization indicated that the most statistically significant ReHo for the 0 to 0.01 Hz band was localized in the DMN regions and subcortical regions, mainly in the thalamus and basal ganglia. For the 0.073 to 0.198 Hz band, the most statistically significant ReHo was mainly localized in the posterior part of the DMN. Finally, for the 0.198 to 0.25 Hz band, the most statistically significant ReHo was localized in the posterior portion of the precuneus/posterior cingulate cortex, with a lower level of local synchronization than typical low frequency bands.

Altered Frequency-Specific Local Synchronization in IGE

The 2-way repeated-measures ANOVA of local synchronization showed a significant main effect of group (Figure 2, axial maps) and a significant main effect of frequency (data not shown). To localize the brain regions that exhibited alterations in local synchronization in IGE patients, we performed separate voxelwise post hoc 3-sample t tests within group main effect mask for each of the 5 frequency bands (Figure 2, inflated surface maps). The spatial extent of brain regions showing group differences gradually expanded when the frequency ranges increased. The BOLD signal of 5 frequency bands, especially the lower bands (0–0.01, 0.01–0.027, and 0.027–
0.073 Hz), showed significantly decreased local synchronization between the 2 groups. IGE patients showed decreased local BOLD oscillation synchronization in lower frequency bands, primarily in the bilateral superior and inferior frontal gyrus, superior and inferior parietal gyrus, and inferior temporal gyrus. These results were consistent with our previous ReHo studies. Only the higher frequency bands (0.073–0.198 and 0.198–0.25 Hz) showed significantly increased local synchronization between the 2 groups, especially in the pregenual anterior cingulate cortex (pgACC), dorsal ACC (dACC), and bilateral insular cortices (see Table 2).

**Frequency-Specific Alternating of Local Synchronization Pattern**

The 2-way repeated-measures ANOVA revealed a significant “frequency by group” interaction effect (Figure 3). The affected brain regions mainly included the anterior and posterior parts of the DMN and subcortical regions, as listed in Table 3. Furthermore, this finding was largely reproduced when using a scrubbing approach to censor “bad” volumes (S-Figure 1, http://links.lww.com/MD/A381). The ROI analysis showed 2 trajectories for “frequency by group” interaction effects (Figure 4). The posterior part of the DMN, insular cortex,
and subcortical regions (first 2 rows) showed a trajectory wherein local synchronization increased with frequency more strongly in the IGE compared with the NC. Conversely, the anterior parts of the DMN (bottom row) showed a monotone decrease in local synchronization.

High-Frequency Local Synchronization Associated With Disease Severity

For the clinical relevance analysis, we found a significant correlation in the right middle cingulate cortex at higher frequency bands. The local synchronization (mean ReHo value) of the right middle cingulate cortex positively correlated with seizure frequency ($r = 0.24$, $P = 0.03$) at the 0.198 to 0.25 Hz band (Figure 5). We found no significant correlation between the local synchronization and the disease duration, onset age, or seizure frequency for the remaining frequency bands.

| Brain Region | BA | MNI (X, Y, Z) | Peak t Value | Cluster Size, mm$^3$ |
|--------------|----|--------------|--------------|---------------------|
| Occipital lobe, R | 19 | 27/81/9 | 3.56 | 729 |
| Occipital lobe, L | 19 | 30/69/21 | 3.45 | 5022 |
| Temporal lobe, R | 39 | 39/48/27 | 3.44 | 594 |
| Rolandic, R | 48 | 48/0/18 | 4.32 | 1998 |
| Rolandic, L | 48 | 48/3/18 | 4.03 | 1080 |
| Caudate, R | N/A | 18/18/15 | 4.53 | 2592 |
| Caudate, L | N/A | 12/0/21 | 3.74 | 1188 |
| Insula, L | 48 | 42/15/21 | 4.19 | 2997 |
| Insula, R | 48 | 39/33/24 | 3.81 | 1188 |
| Pregenual anterior cingulate, R | 32 | 6/33/0 | 4.33 | 1188 |
| Dorsal anterior cingulate, R | 24 | 3/0/21 | 3.48 | 594 |
| Occipital lobe, R | 19 | 30/61/21 | 3.56 | 2997 |
| Occipital lobe, L | 19 | 30/81/0 | 3.02 | 1053 |
| Parietal lobe, R | N/A | 12/60/33 | 3.67 | 729 |
| Parietal lobe, L | N/A | 15/57/42 | 4.09 | 1350 |
| Middle orbitofrontal gyrus, R | 47 | 30/48/9 | 4.17 | 1836 |
| Inferior frontal gyrus, L | 48 | 33/30/9 | 3.94 | 1188 |
| Frontal lobe, R | 11 | 18/48/6 | 4.30 | 2970 |
| Superior temporal gyrus, R | 22 | 51/30/9 | 3.34 | 594 |
| Caudate, L | N/A | 12/15/15 | 4.56 | 1296 |
| Caudate, R | N/A | 15/15/15 | 4.99 | 2349 |
| Insula, R | 48 | 36/18/12 | 3.83 | 810 |
| Insula, L | 48 | 33/3/18 | 4.70 | 1809 |
| Anterior cingulate, R | 32 | 3/42/9 | 3.55 | 945 |
| Cerebelum 8, R | N/A | 21/66/42 | 3.86 | 1728 |
| Cerebelum crus1, L | N/A | 48/69/27 | 3.49 | 540 |
| Middle frontal gyrus, R | 11 | 30/48/12 | 4.70 | 2597 |
| Frontal lobe, L | 47 | 36/30/6 | 4.02 | 648 |
| Superior frontal gyrus, R | 32 | 15/30/42 | 4.39 | 2781 |
| Middle temporal gyrus, R | 37 | 42/66/6 | 3.92 | 648 |
| Occipital lobe, L | 37 | 30/57/6 | 3.79 | 621 |
| Parietal lobe, R | 4 | 57/18/51 | 4.17 | 540 |

BA = Brodmann area, MNI = Montreal Neurological Institute, L = left hemisphere, N/A = not available, R = right hemisphere.

DISCUSSION

The previous ReHo analysis at conventional low frequency band would be used to detect intrinsic brain activity in IGE patients, which runs the risk of overlooking meaningful information associated with frequency-specific spontaneous BOLD fluctuation. The current study is the first to specifically investigate frequency-specific alterations of BOLD local synchronization in patients with IGE. First, IGE patients showed decreased local synchronization primarily in the DMN, and predominately at lower frequency bands. However, IGE patients also showed increased frequency-specific local synchronization in the pgACC, dACC, and bilateral insular cortices at higher frequencies. Additionally, we observed a frequency-specific alternation pattern in the DMN such that differences in local synchronization increased with the frequency range. Importantly, the aberrant high-frequency local synchronization in the middle cingulate cortex was associated with disease...
duration, thus linking BOLD frequency changes to disease severity. Taken together, these findings provide an overall view of frequency-specific local synchronization of BOLD oscillations, and may be useful in uncovering abnormal synchronous neuronal activity in IGE at specific frequency bands.

Full Bandwidth Analysis of Local Synchronization Features

Recent studies have examined resting-state fMRI for higher-frequency BOLD oscillations and functional connectivity in the diseased brain.22,23,30 The present work is a natural extension of our previous resting-state fMRI study, which detected BOLD local synchronization in patients with IGE at low frequencies only (0.01–0.08 Hz).18 Here, we offer some new insights regarding frequency-specific local synchronization spanning the full bandwidth, which agree with and expand upon the findings of a recent report showing varied ReHo across 2 bands.31

Altered Frequency-Specific Local Synchronization in IGE

It is not surprising that we observed altered local synchronization of BOLD fluctuations, primarily in the DMN, in patients with IGE, as this is consistent with the previous findings.18 The resting-state activity of this network seems to be nonspecific, not only because of transient-seizure-suspended DMN activation in various phenotypes of IGE,28,32,33 but also because of long-term functional impairments of the DMN in focal epilepsy.37,34 Interestingly, we found decreased local synchronization predominately in the lower frequency bands (0–0.01, 0.01–0.027, and 0.027–0.073 Hz) (Figure 2). This suggests that frequency-specific functional DMN impairment in patients with IGE is dominated by lower frequency BOLD fluctuations.

Our most remarkable finding was that patients with IGE exhibit significantly increased local synchronization in the pgACC, dACC, and bilateral insular cortices in higher frequency bands (Figure 2). These regions are components of a "conscious perception network,"35 which is associated with conscious perception by synchronized oscillations.36,37 To the best of our knowledge, this finding has for the first time been reported in a human functional neuroimaging study of epilepsy. Generalized tonic–clonic seizures, also named grand mal seizures, are a dramatic type of convulsive seizure. After these convulsive episodes, consciousness is deeply impaired.37 Cerebral blood flow studies have an advantage in that they employ an injection of radiotracer during a seizure attempt. This enables researchers to identify abnormal activity that possibly contributes to the ictal impairment of consciousness.38–40 Our findings support the notion that aberrant local synchronization occurs in the “conscious perception network" in patients with IGE. Note that the pattern of between-group differences that we observed was only found in higher frequency bands, suggesting a frequency-specific disruption. More recently, epilepsy-related high-frequency oscillations in electroencephalography have
captured pathological activities, such that they can now be considered new biomarkers of the extent and intensity of epileptogenicity. Although the correspondence between the BOLD frequency bands investigated here and EEG rhythms is not well known, it seems possible that the higher frequency band of BOLD fluctuations may play a key role in future clinical therapy. Some therapeutic tools, such as electroconvulsive therapy, have the advantage of controlling timing and producing a relatively consistent seizure onset in an experimental setting. However, the exact mechanism of action remains unknown. Some studies have suggested that the resting-state functional connectivity pattern may correspond with the hierarchical functions of the DMN, indicating that the DMN is modulated by frequency in patients with IGE. This finding leads to a novel pathophysiological perspective: the hierarchical functions of the DMN may correspond with the frequency-specific local synchronization of BOLD fluctuations. Taken together, these results demonstrate that anterior and posterior DMN are implicated in disease and frequency bands, thus extending our understanding of frequency-specific pathology in patients with IGE.

**TABLE 3. Brain Regions Showing Statistically Significant “Frequency by Group” Interaction Effects**

| Brain Region                     | BA     | MNI (X, Y, Z) | Peak F–z Value | Cluster Size, mm³ |
|----------------------------------|--------|--------------|----------------|-------------------|
| Thalamus, L                      | N/A    | −18 −18 15   | 3.04           | 594               |
| Thalamus, R                      | N/A    | −15 −12 18   | 2.95           | 675               |
| Insula, L                        | 13     | −33 21 0     | 2.95           | 2133              |
| Insula, R                        | 13     | 33 24 0      | 3.12           | 621               |
| Supramarginal gyrus, L           | 40     | −57 −30 33   | 2.97           | 1566              |
| Supramarginal gyrus, R           | 40     | 60 −21 27   | 2.84           | 2619              |
| Middle cingulate cortex          | 24     | 0 −12 36    | 3.26           | 1134              |
| Middle cingulate cortex R        | 32     | 9 15 36     | 2.77           | 945               |
| Caudate, L                       | N/A    | −15 3 15    | 2.38           | 594               |
| Precuneus, R                     | 31/7   | 15 −66 30   | 3.15           | 1323              |
| Middle orbital frontal gyrus, L  | 10/11  | −30 57 −9   | 3.30           | 1539              |
| Medial superior frontal gyrus, R | 9      | 3 51 27     | 2.61           | 594               |
| Middle temporal gyrus, L         | 20     | −45 −6 −39  | 4.65           | 4833              |
| Middle temporal gyrus, R         | 20     | 36 −15 −36  | 3.23           | 567               |
| Cerebellar tonsil, R             | N/A    | 3 −51 −42    | 3.68           | 1593              |
| Cerebellum crus 2, L             | N/A    | −45 −75 −39 | 4.34           | 5049              |
| Cerebellum crus 1, L             | N/A    | −45 −45 −39 | 4.13           | 1593              |
| Declive, R                       | N/A    | 18 −63 −21  | 3.46           | 1053              |
| Superior temporal gyrus, R       | 38     | 60 18 −24   | 2.98           | 864               |
| Middle occipital gyrus, L        | 18/19  | −33 −96 9   | 5.18           | 4752              |
| Superior temporal gyrus, L       | 22     | −48 −54 −3  | 3.10           | 1755              |
| Inferior frontal gyrus, R        | 46     | 51 39 9     | 2.79           | 702               |
| Middle occipital gyrus, R        | 19     | 36 −87 33   | 4.36           | 5400              |
| Parietal lobule, R               | 7/19   | 9 −81 54    | 3.92           | 8235              |
| Paracentral lobule, L            | 6      | −12 −18 72  | 3.25           | 1890              |

BA = Brodmann area, MNI = Montreal Neurological Institute, L = left hemisphere, N/A = not available, R = right hemisphere.

**Frequency Alternating of Local Synchronization Pattern in IGE**

Across 5 frequency bands, we observed 2 distinct trajectories of shift in different networks, along with an increasing frequency range (Figure 4). This suggests a frequency-specific alternating feature of local synchronization in patients with IGE. The posterior portion of the DMN and subcortical regions showed increased local synchronization as the frequency range increased. Conversely, the anterior portion of the DMN showed a local synchronization shift that decreased in a monotone way. On the one hand, altered local synchronization in both anterior and posterior portions of the DMN reflects the long-term injurious effects of epileptic activity, indicating defective inhibition of the DMN in IGE. This finding is consistent with the previous EEG-fMRI and functional connectivity studies. On the other hand, the gradual reversal of local synchronization across 5 frequency bands between 2 subnetworks of the DMN indicates that the DMN is modulated by frequency in patients with IGE. Note that the DMN may encompass frequency-specific anatomy-specific subcomponents, which is associated with the hierarchical functions of the default mode. This finding leads to a novel pathophysiological perspective: the hierarchical functions of the DMN may correspond with the frequency-specific local synchronization of BOLD fluctuations. Taken together, these results demonstrate that anterior and posterior DMN are implicated in disease and frequency bands, thus extending our understanding of frequency-specific pathology in patients with IGE.

**Limitations**

There are several limitations of this study. First, 3 IGE subgroups used the current study. Previous studies have suggested that the resting-state functional connectivity pattern varies among IGE subcategories. However, this study did not simultaneously acquire scalp EEG, which would have enabled us to exclude MRI data with GSWDs. Further simultaneous EEG-MRI data would enable us to examine the specific relationship between local frequency-specific synchronization and interictal epileptic discharges. Third, we did not consider the socioeconomic status for the patients and controls; however, it is unlikely that these factors contributed to the between-groups difference because most of the participants were from middle-class families.
Finally, the local synchronization was measured at a low sampling rate (Repetition Time = 2 seconds), which impeded investigations of high-frequency alternation. Future research could use advanced data acquisition sequences to enable whole-brain fMRI scanning at a subsecond temporal resolution.

CONCLUSIONS

The present study examined the frequency-specific local synchronization of BOLD fluctuations during the resting state, taking into account the full bandwidth. First, IGE patients showed decreased local synchronization in low frequency, primarily in the DMN. We also found increased local synchronization in high frequency in the “conscious perception network,” possibly contributing to the impairment of consciousness. Additionally, we found frequency-specific alternation of local synchronization in the posterior portion of the DMN, suggesting an interaction between epilepsy and frequency bands. Importantly, the aberrant higher-frequency local synchronization in the middle cingulate cortex was associated with disease duration, thus linking BOLD frequency-specific changes to disease severity. In general, these findings provide an overall view of frequency-specific local synchronization of BOLD fluctuations, and may have potential in uncovering abnormal synchronous neuronal activity at specific frequency bands in patients with IGE.

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