Time-Shift Homotopic Connectivity in Mesial Temporal Lobe Epilepsy


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ABSTRACT

BACKGROUND AND PURPOSE: Voxel-mirrored intrinsic functional connectivity allows the depiction of interhemispheric homotopic connections in the human brain, whereas time-shift intrinsic functional connectivity allows the detection of the extent of brain injury by measuring hemodynamic properties. We combined time-shift voxel-mirrored homotopic connectivity analyses to investigate the alterations in homotopic connectivity in mesial temporal lobe epilepsy and assessed the value of applying this approach to epilepsy lateralization and the prediction of surgical outcomes in mesial temporal lobe epilepsy.

MATERIALS AND METHODS: Resting-state functional MR imaging data were acquired from patients with unilateral mesial temporal lobe epilepsy (n = 62) (31 left- and 31 right-side) and healthy controls (n = 33). Dynamic interhemispheric homotopic architecture seeding from each hemisphere was individually calculated by 0, 1, 2, and 3 repetition time-shift voxel-mirrored homotopic connectivity. Voxel-mirrored homotopic connectivity maps were compared between the patient and control groups by using 1-way ANOVA for each time-shift condition, separately. Group comparisons were further performed on the laterality of voxel-mirrored homotopic connectivity in each time-shift condition. Finally, we correlated the interhemispheric homotopic connection to the surgical outcomes in a portion of the patients (n = 20).

RESULTS: The patients with mesial temporal lobe epilepsy showed decreased homotopic connectivity in the mesial temporal structures, temporal pole, and striatum. Alterations of the bihemispheric homotopic connectivity were lateralized along with delays in the time-shift in mesial temporal lobe epilepsy. The patients with unsuccessful surgical outcomes presented larger interhemispheric voxel-mirrored homotopic connectivity differences.

CONCLUSIONS: This study showed whole patterns of dynamic alterations of interhemispheric homotopic connectivity in mesial temporal lobe epilepsy, extending the knowledge of abnormalities in interhemispheric connectivity in this condition. Time-shift voxel-mirrored homotopic connectivity has the potential for lateralization of unilateral mesial temporal lobe epilepsy and may have the capability of predicting surgical outcomes in this condition.

ABBREVIATIONS: HC = healthy controls; mTLE = mesial temporal lobe epilepsy; ts = time-shift; VMHC = voxel-mirrored homotopic connectivity
increases in the higher order processing areas, which provided insight into the evolution of brain development. Studies have also revealed specific alterations of homotopic connectivity in a cohort of brain diseases.6-8 Decreased VMHC in schizophrenia has been suggested to reflect the substantial impairment of interhemispheric coordination in these patients.6 Anderson et al9 found homotopic connectivity alterations related to behavioral and developmental abnormalities in autism.7 More recently, studies have further correlated functional homotopic connectivity with microstructural impairment in multiple sclerosis8 and idiopathic generalized epilepsy.9

In contrast to the brain disorders featuring abnormal connection pathways as mentioned above,6,8 mesial temporal lobe epilepsy (mTLE) is a location-related disease characterized by hippocampal sclerosis.10 Unilateral mTLE can cause bilateral and distributed brain impairments due to seizure propagation via the mesial temporal epileptic network.11-13 Resting-state fMRI studies have shown asymmetric connections between hemispheres14-16 and decreased connectivity between bilateral hippocampi in mTLE.17,18 These findings suggest that there are intrahemispheric connection abnormalities in unilateral mTLE. However, neither the homotopic alterations of whole-brain functional connectivity nor the relationship between asymmetric lesions and interhemispheric communication in this disease has been investigated.

fMRI-based VMHC provides a feasible way to observe the whole-brain homotopic connectivity alterations in mTLE. However, the conventional nondirectional functional connectivity measure, as used in VMHC, cannot detect the connection abnormalities resulting from deficits of the seed or target region. Recently, Lv et al19 proposed a time-shift (ts) analysis for resting-state functional connectivity. They quantified the temporal shift correlation between time courses of each voxel and global mean signal19 and correlated the time shifts with the extent and degree of perfusion delay in patients with stroke.19,20 In addition, our previous study used time-shift correlation analysis to demonstrate the sequential effects of epileptic discharges on intrinsic network connectivity in children with absence epilepsy.21 Thus, time-shift delays in resting-state spontaneous connectivity were assumed to reflect brain hemodynamics and could measure the degree of brain injury; time-shift analysis also provides directional information as a measure of functional connectivity.21 In the current work, we combined the time-shift connectivity with the VMHC technique and applied them to resting-state fMRI data from patients with unilateral mTLE. We hypothesized that this strategy would allow us to assess the whole-brain homotopic connection impairments resulting from different hemispheres and may potentially be a tool for epileptic focus lateralization and surgical outcome prediction in mTLE.

MATERIALS AND METHODS

Subjects
Sixty-two patients with unilateral mTLE (31 left-sided and 31 right-sided) were recruited (demographic and clinical information are detailed in the Table). Part of the patient population (23 patients with left-sided mTLE) was reported in our previous work in which the Granger causal information of the hippocampus in left-sided mTLE was estimated.13 Diagnosis of mTLE was performed according to the International League Against Epilepsy 2001 classification based on a comprehensive evaluation, including seizure history and semiology, neurologic examination, diagnostic MR imaging, and electroencephalography records in all patients. Patients were selected on the basis of several criteria: 1) Patients who were young and middle-aged were included (patients younger than 18 years or older than 50 years were excluded); 2) patients who exhibited identifiable structural MR imaging abnormalities other than hippocampal sclerosis, such as cortical dysplasia, vascular malformation, or tumor were excluded. Twenty patients of the 62 underwent standard anterior temporal lobectomy: Fourteen cases (8 left- and 6 right-sided mTLE) had successful surgical outcomes, and 6 cases (4 left- and 2 right-sided mTLE) had unsuccessful outcomes. A successful outcome was defined as seizure-free with or without auras at the latest follow-up that occurred at least 1 year after the surgery, which satisfies the International League Against Epilepsy Outcome Criteria 1 or 22 and Engel category 1A or 1B.23 No postoperative MR imaging data were used in this article.

Moreover, healthy controls (HC) (n = 33) were recruited from the hospital staff. These controls did not have neurologic or psychiatric disorders at the time of the study. There were no differences in ages and sexes between the controls and patients (Table).

This study was approved by the local Medical Ethics Committee, and written informed consent was obtained from all the participants.

Data Acquisition
We acquired functional and structural images in patients and HC by using a Trio 3T scanner (Siemens, Erlangen, Germany). We used foam padding to minimize head motion. We acquired resting-state functional images by using a single-shot, gradient-recalled echo-planar imaging sequence (250 volumes, TR = 2000 ms, TE = 30 ms, flip angle = 90°, FOV = 240 × 240 mm²,

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>L-mTLE (n = 31)</th>
<th>R-mTLE (n = 31)</th>
<th>HC (n = 33)</th>
<th>ANOVA (P Value)</th>
<th>L-mTLE vs R-mTLE (P Value)</th>
<th>L-mTLE vs HC (P Value)</th>
<th>R-mTLE vs HC (P Value)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yr)</td>
<td>26.5 ± 8.4</td>
<td>29.7 ± 8.5</td>
<td>26.7 ± 7.3</td>
<td>0.223^a</td>
<td>0.146^b</td>
<td>0.903^c</td>
<td>0.344^c</td>
</tr>
<tr>
<td>Sex</td>
<td>19 M/12 F</td>
<td>16 M/15 F</td>
<td>21 M/12 F</td>
<td>0.592^b</td>
<td>0.442^d</td>
<td>0.846^d</td>
<td>0.330^d</td>
</tr>
<tr>
<td>Successful outcome</td>
<td>8</td>
<td>6</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Unsuccessful outcome</td>
<td>4</td>
<td>2</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Note: N/A indicates not applicable; L, left; R, right.

^a One-way analysis of variance.
^b Kruskal-Wallis ANOVA.
^c Two-sample t test.
^d χ² test.
intersection gap = 0.4 mm, voxel size = 3.75 × 3.75 × 4 mm³, 30 transverse sections aligned along the anterior/posterior commissure). Subjects were instructed simply to rest with their eyes closed, not to think about anything in particular, and not to fall asleep. Subsequently, we acquired high-resolution 3D T1-weighted anatomic images in a sagittal orientation by using a magnetization-prepared rapid acquisition of gradient echo sequence (TR = 2300 ms, TE = 2.98 ms, flip angle = 9°, FOV = 256 × 256 mm², voxel size = 1 × 1 mm³, 176 sections without intersection gap).

Data Preprocessing
Functional image preprocessing was performed by using the DPARSF 2.2 (http://www.restfmri.net) and SPM8 (Wellcome Department of Imaging Neuroscience, London, UK; http://www.fil.ion.ucl.ac.uk/spm) toolkits. We excluded the first 10 images to ensure steady-state longitudinal magnetization, and then we corrected the remaining images for temporal differences and head motion. No translation or rotation parameters in any given dataset exceeded ±1 mm or ±1°. We then coregistered individual 3D T1 images to functional images. The 3D T1-weighted anatomic images were segmented into gray matter, white matter, and CSF by using unified segmentation. Then, a nonlinear spatial deformation was calculated from the gray matter images to a gray matter template.

FIG 1. Workflow for time-shift voxel-mirrored homotopic connectivity calculation. A, Ts-VMHC of seeding from the left hemisphere. The left side was set as the seeding side; the right side, as the targeting side. The 0 TR, 1 TR, 2 TR, and 3 TR shifted time-courses of each voxel on the targeting side (blue line) were separately correlated with the time course of the mirrored voxel on the seeding side (red line), which generated 0 TR, 1 TR, 2 TR, and 3 TR ts-VMHC hemispheric maps of the seeding (left) side. B, Similarly, 4 hemispheric ts-VMHC maps of the right side were also generated when taking the right hemisphere as the seeding side and the left hemisphere as the targeting side.

FIG 2. Group comparisons of ts-VMHC maps between the patients and healthy controls. The left and right hemispheric ts-VMHC maps (0 TR, 1 TR, 2 TR, and 3 TR) of each patient group were compared with those of the HC by using 2-sample t tests ($P < .05$, AlphaSim corrected) under ANOVA. For a better visualization, we combined the comparison results of the left and right hemispheric ts-VMHC maps with a whole-brain map. The mesial temporal structures, limbic/paralimbic areas, and striatum areas show time-shift-dependent VMHC decreases.
matter template in Montreal Neurological Institute space. This transformation was then applied to the functional images, which were resectioned at a resolution of 3 × 10 mm3. The data were bandpass-filtered (0.01–0.08 Hz). Linear regression was performed to remove the effect of 6 head-motion-averaged signals from the CSF, white matter, and the global brain signal.25 The residual images were further coregistered to a symmetric template according to previous studies.5,8 The symmetric template was created as follows: First, all participants’ normalized gray matter images were averaged to create a mean normalized gray matter template. This template was then averaged with its left-right mirrored version to generate a group-specific symmetric template.5 Finally, we spatially smoothed the images with an 8-mm full width at half maximum isotropic Gaussian kernel.

**Time-Shift Voxel-Mirrored Homotopic Connectivity**

We combined the time-shift analysis19 with voxel-mirrored homotopic connectivity analysis.3 We first took the left hemisphere as the seeding side and the right hemisphere as the targeting side. The 0 TR, 1 TR, 2 TR, and 3 TR shifted time-courses of each voxel on the targeting side were separately correlated with the time course of the mirrored voxel on the seeding side, which generated 0 TR, 1 TR, 2 TR, and 3 TR ts-VMHC hemispheric maps of the seeding (left) side. Similarly, 4 hemispheric ts-VMHC maps of the right side were also generated when taking the right hemisphere as the seeding side and the left hemisphere as the targeting side (Fig 1). The correlation maps were then converted to z scores by using the Fisher r-to-z transformation.

**Group Comparisons of ts-VMHC**

In the second-level analyses, the left and right hemispheric ts-VMHC maps (0 TR, 1 TR, 2 TR, and 3 TR) of each patient group were compared with those of the HC by using a 2-sample t test (AlphaSim corrected P < .05, combined height threshold P < .01, and extent threshold k > 20 voxels). For a better visualization, we combined the comparison results of the left and right hemispheric ts-VMHC maps with a whole-brain map (Fig 2 and On-line Table).

**Laterality Analyses of ts-VMHC**

To quantitatively identify the difference between the left and right hemispheric ts-VMHC and to further determine whether the ts-VMHC could be used for mTLE lateralization, we performed a voxelwise laterality analysis. In each individual of all 3 participant groups, the right hemispheric ts-VMHC map was subtracted from the left hemispheric ts-VMHC map and the results were divided by the summation of both maps. Then the laterality (hemispheric) maps of all 3 groups of participants were compared by using 1-way ANOVA implemented in SPM8 in each time-shift condition (P < .05, AlphaSim correction). Then, a post hoc region of interest–based analysis was subsequently performed. The ROIs were selected from the above ANOVA result in 3 TR time-shift conditions. The laterality values in the ROIs were extracted for ANOVA among the 3 participant groups (Fig 3).

**Predicting Surgical Outcomes of ts-VMHC**

We performed a preliminary analysis to retrospectively investigate the relationship between ts-VMHC and surgical outcomes of mTLE. Twenty patients who underwent anterior temporal lobectomy were included. The patients with right-sided mTLE (R-mTLE) were left-right flipped to produce a homogeneous left-sided mTLE dataset. We calculated the difference of VMHC between bihemispheres by subtracting the connectivity values of the contralateral side from those of the lesional side (ie, the epileptogenic side). Subsequently, the interhemispheric differences were compared by using 2-sample t tests between the patients with successful surgical outcomes and those with unsuccessful outcomes.

**RESULTS**

**Group Comparison Results of ts-VMHC**

On the basis of the comparison of the results of VMHC without time-shift (ie, 0 TR shift), both patient groups showed decreased homotopic connectivity in the regions of limbic/paralimbic areas (mainly in the mesial temporal structures including the hippocampus/parahippocampus, temporal pole, amygdala, and insula) and striatum areas (mainly in the putamen and pallidum) relative to the HC (P < .05 AlphaSim correction). Along with the
and Fig 2).

Connectivity was detected in the patient groups (On-line Table 1750). First, our results showed connectivity analysis with VMHC analysis and applied them to time-shift functional connectivity. In this study, we innovatively combined time-shift functional connectivity with VMHC analysis and applied them to the mesial temporal structures, temporal pole, amygdala, insula, putamen, and pallidum. These regions have been regarded as key nodes in the mesial temporal epileptic networks and play important roles in the generation, propagation, and modulation of epileptic activity. Imaging studies have demonstrated structural and functional abnormalities in these regions. Thus, we considered that reduced homotopic connectivity might result from nodal impairments of these regions in mTLE. Moreover, studies have proposed a variety of propagation patterns of epileptic activity from the originating focus to the contralateral side; the limbic structures and basal ganglia are the important nodes within the propagation pathway. Here we cautiously proposed another explanation, that time-shift-dependent alterations of VMHC of these regions might represent propagations of epileptic activity.

Along with the time-shift delays, the decreases in homotopic connectivity in both hemispheres tended to be weaker, whereas the decreases on the lesioned side (ie, the left side in the left-sided mTLE and the right side in the right-sided mTLE patients) were more persistent. Until the 3 TR shift, only the difference in the mesial temporal structures of the lesioned side survived. No increase in homotopic connectivity was detected in the patient groups (On-line Table and Fig 2).

**Laterality Analyses of ts-VMHC**

Along with the time-shift delays at 1 TR, 2 TR, and 3 TR, the results of the 1-way ANOVA among subject groups showed more significant laterality of homotopic connectivity in the mesial temporal structures. Post hoc region-of-interest analyses repeated the above 3 TR time-shifted ANOVA results (Fig 3). Along with time shifts of 1 TR, 2 TR, and 3 TR, the difference between the patients with left-sided mTLE and those with right-sided mTLE became more significant ($P = .001$ in 1 TR, $P < .001$ in 2 TR, and $P < .001$ in 3 TR).

**Predicting Surgical Outcomes**

The preliminary analysis correlating ts-VMHC with surgical outcomes showed that the patients with unsuccessful surgical outcomes had larger VMHC differences between the hemispheres than those with successful outcomes at each time-shift condition (1 TR-shifted, $P = .001$; 2 TR-shifted, $P = .001$; 3 TR-shifted, $P = .008$). Moreover, along with time shifts of 1 TR, 2 TR, and 3 TR, the interhemispheric VMHC differences increased in both the groups with successful and unsuccessful outcomes (Fig 4).

**DISCUSSION**

In this study, we innovatively combined time-shift functional connectivity analysis with VMHC analysis and applied them to data from patients with unilateral mTLE. First, our results showed decreased homotopic connectivity in the mesial temporal structures, temporal pole, and striatum in mTLE, which provided the first complete picture of interhemispheric connectivity abnormalities in mTLE. Second, we found that the alterations in the bihemispheric homotopic connectivity led to lateralization along with time-shift delays in patients with mTLE. This finding suggests that ts-VMHC may have the potential for lateralization of unilateral mTLE. Third, the hemispheric differences in homotopic connectivity were related to surgical outcomes in patients with mTLE, which indicates that the time-shift homotopic connectivity may be predictive of surgical outcomes in mTLE.

We first revealed the whole-brain patterns of altered homotopic connectivity in mTLE. We found that the mesial temporal structures, including the hippocampus and parahippocampal gyrus, showed decreased homotopic connectivity, consistent with previous studies. Furthermore, we found decreased homotopic connectivity in the temporal pole, amygdala, insula, putamen, and pallidum. These regions have been regarded as key nodes in the mesial temporal epileptic networks and play important roles in the generation, propagation, and modulation of epileptic activity. Imaging studies have demonstrated structural and functional abnormalities in these regions. Thus, we considered that reduced homotopic connectivity might result from nodal impairments of these regions in mTLE. Moreover, studies have proposed a variety of propagation patterns of epileptic activity from the originating focus to the contralateral side; the limbic structures and basal ganglia are the important nodes within the propagation pathway. Here we cautiously proposed another explanation, that time-shift-dependent alterations of VMHC of these regions might represent propagations of epileptic activity.

Along with the time-shift delays, the decreases of the bihemispheric homotopic connectivity were more lateralized in patients with unilateral mTLE. The lesioned side showed more persistent decrease in homotopic connectivity. We introduced a time-shift functional connectivity analysis, different from the conventional VMHC approach, which provided directional information to identify connectivity abnormalities resulting from different nodal impairments. Time-shift functional connectivity was first proposed by Lv et al in their fMRI study on stroke. They proposed that the time-shift delays in resting-state fMRI connectivity represent the hemodynamics/perfusion property of stroke and could be used to measure the extent of the brain lesions. Here, we assume that different decreases in the time-shift homotopic connectivity between the hemispheres may result from different hemodynamic properties between the hemispheres in mTLE. Previous studies showed that the epileptic focus presented interictal hypoperfusion in mTLE. This finding confirmed our hypothesis that the strategy of combining time-shift functional connectivity with VMHC had the potential to lateralize unilateral mTLE.
Our preliminary analysis showed that the results of ts-VMHC could be related to the surgical outcomes in patients with mTLE. The patients with unsuccessful outcomes had more interhemispheric VMHC differences. The finding suggested that the patients with unsuccessful outcomes might have more severe lesions and presented more disruption of interhemispheric communication. Previous studies have shown that resting-state fMRI has the capability of predicting surgical outcomes in mTLE.16,38 Different from those studies that used predefined region-of-interest analysis,16,38 our study used an unbiased voxelwise analysis approach and simultaneously observed the dynamic property of intrinsic connectivity in mTLE. Our study provided more comprehensive information for resting-state fMRI in the prediction of surgical outcome in mTLE.

**Limitations**

A few of the limitations in this study are noteworthy. First, we did not address the possible effects of antiepileptic drugs on fMRI in the study patients. Second, another presumption of time-shift connectivity might reflect propagation of epileptic activation; simultaneous electroencephalography and fMRI to record interictal epileptic discharges are needed in future studies. Third, the included patients had detectable hippocampal sclerosis, which might weaken the significance of epilepsy lateralization. Future study on patients with negative MR imaging findings is needed to validate this study. Finally, this work only observed the interhemispheric homotopic connectivity; measures of intrahemispheric and whole-brain connectivity might benefit from the understanding of the underlying mechanisms of connectivity alterations in epilepsy.

**CONCLUSIONS**

This study combined time-shift functional connectivity analysis with VMHC analysis and observed ts-VMHC alteration in patients with mTLE. The results showed the whole-brain pattern of aberrant interhemispheric connectivity in mTLE and further implicated the associated nodal impairment. Moreover, our study provided a novel fMRI approach that has potential value for epilepsy.

**REFERENCES**