Altered interhemispheric functional connectivity in remitted bipolar disorder: A Resting State fMRI Study

Lianping Zhao¹,², Ying Wang¹,³, Yanbin Jia⁴, Shuming Zhong⁴, Yao Sun³, Zhangzhang Qi¹, Zhongping Zhang⁵ & Li Huang¹

Abnormalities in structural and functional brain connectivity have been increasingly reported in patients with bipolar disorder (BD). However, alterations of remitted BD (RBD) in functional connectivity between the cerebral hemispheres are still not well understood. This study was designed to analyze the pattern of the interhemispheric functional connectivity of the whole brain in patients with remitted BD during resting state. Twenty patients with RBD and 38 healthy controls (HC) underwent the resting-state functional magnetic resonance imaging. The functional connectivity between any pair of symmetrical interhemispheric voxels (i.e., functional homotopy) was measured by voxel-mirrored homotopic connectivity (VMHC). The patients with RBD showed lower VMHC than HC in the middle frontal gyrus and precentral gyrus. No regions of increased VMHC were detected in the RBD patients. There were no significant correlations between the VMHC values in these regions and the clinical variables. These findings suggest substantial impairment of interhemispheric coordination in RBD and they may represent trait, rather than state, neurobiological feature of brain function in BD.

Bipolar disorder (BD) is a chronic, severe, and fluctuating mental disorder and the worldwide prevalence of the BD spectrum is 1–4%¹. The psychiatric condition of BD commonly characterized by cyclically alternating depressive and manic episodes interspersed with symptomatic remissions. Symptomatic remissions are not free of residual cognitive and/or emotional symptoms²,³. Some of these symptoms actually represent trait characteristics of the illness. Although researchers agree that BD is multifactorial with inherited, genetic and environmental risk factors, the neuropathological mechanisms are still remains unclear. The neuropathological changes of remitted BD (RBD) are most close to the pathophysiological determinants in that it may elucidate potential neurobiological mechanisms of BD while mitigating the possible confounding effects of current mood state, illness severity, nonspecific effects of chronic illness and stress. Consequently, identification of neuropathological alterations in RBD may contribute to early diagnosis of BD thus reducing the latency to adequate treatment and improving outcome.

Evidences from literatures have revealed that the imbalance between left and right hemispheric activity may contribute to the pathophysiology of BD⁴–⁷. The corpus callosum (CC), the largest white matter structure connecting the left and right cerebral hemispheres, plays a pivotal role in interhemispheric communication and coordination, especially in the integration of emotional, cognitive, motor, and sensory information. Structural magnetic resonance imaging (MRI) studies have found area and shape changes in the CC in BD⁸. In addition, diffusion tensor imaging (DTI) and tractography studies have found reduced fractional anisotropy (FA) in the genu, body, and splenium of the CC in patients with BD during mood episodes⁷,⁹,¹⁰ and in euthymic phase relative to controls, suggesting that disruption of the organization of fiber tracts may be related to efficiency of interhemispheric signal transfer in the case of the CC. A recent study reported impaired interhemispheric but
relatively preserved intrahemispheric integration in BD by DTI and graph theoretical analysis. However, relatively little is known about the alterations in functional connectivity between the bilateral cerebral hemispheres in patients with RBD.

Resting-state fMRI (rs-fMRI), which reveals the patterns of coherent spontaneous fluctuations of blood oxygen level dependent (BOLD) signals, provides fresh insights into the intrahemispheric functional connectivity. The widely used approaches of rs-fMRI conducted in BD have employed either seed-based or amplitude of low-frequency fluctuation (ALFF) analyses. While informative, the seed-based approach is almost always limited by the inherent bias introduced by predefining regions of interest (the seed region) for connectivity analyses. By contrast, ALFF is a useful tool for identifying group differences in local BOLD signal frequency fluctuations, but it is not a direct measure of functional connectivity. Kenny et al. have also emphasized that affective disorders are more likely due to aberrations at the circuit level rather than at a localized brain region. Functional homotopy is designed to quantify the high degree of synchrony in spontaneous activity between functionally corresponding intrahemispheric counterparts, one of the most salient features of the brain intrinsic functional architecture, likely reflecting the importance of interhemispheric communication to integrated brain function underlying coherent cognition and behavior. Voxel-mirrored homotopic connectivity (VMHC) is a recently validated approach that quantifies the resting-state functional connectivity between each voxel in one hemisphere and its corresponding voxel in the opposite hemisphere. Recent work has demonstrated that VMHC is a reliable and reproducible rs-fMRI metric. Since VMHC first introduced by Zuo et al., it has revealed aberrant interhemispheric functional connectivity in several neuropsychiatric disorders including schizophrenia, cocaine addiction, migraine, somatization disorder, post-traumatic stress disorder and major depressive disorder. Our previous study revealed reduced interhemispheric resting-state functional connectivity in the medial prefrontal cortex and inferior temporal gyrus in unmedicated BD II depression. However, there are no studies to investigate possible impairment of homotopic resting-state functional connectivity in RBD patients so far.

In the present study, we compared the interhemispheric resting-state functional connectivity in patients with RBD and healthy controls (HC) by use of VMHC analysis. Reduced VMHC values in patients with RBD were anticipated, which would suggest an impairment of interhemispheric functional coordination. We believe such data will contribute to our improved understanding of the pathophysiology of BD.

Material and Methods

Subjects. A cohort of 24 right-handed, out- or in-patients with RBD (23 remitted from BD-II, 1 remitted from BD-I) were recruited from the psychiatry department of First Affiliated Hospital of Jinan University, Guangzhou, China. The patients were aged from 18 to 47 years. All patients met DSM-IV criteria for BD-I or II in past diagnosis according to the diagnostic assessment by the Structured Clinical Interview for DSM-IV Patient Edition (SCID-P). At the time of the measurements all patients were in disease remission. Remitted state was defined by Hamilton Rating Scale for Depression-24 (HAM-D-24) scores of <8 and Young Mania Rating Scale (YMRS) scores of <6, and the depressive or manic symptoms disappeared for at least or more than 6 months. At the time of the study, 4 patients in the BD group were without medication and 20 were receiving antidepressants (duloxetine or paroxetine), and/or mood stabilizers (lithium, sodium valproate), and/or atypical antipsychotic medications (olanzapine, risperidone or quetiapine). Exclusion criteria included the presence of (i) any current psychiatric illness (with the exception of BD and anxiety disorders), (ii) a history of electroconvulsive therapy, (iii) any history of moderate/severe head injury, head trauma, neurological disorder, or mental retardation, (iv) alcohol/substance abuse or dependence, (v) the presence of a concurrent significant physical illness, and (vi) pregnancy or any contraindication to MRI scanning. A total of 42 right-handed age and gender matched healthy control subjects were recruited via local advertisements. They were carefully screened through a diagnostic interview, the Structured Clinical Interview for DSM-IV Nonpatient Edition (SCID-NP), to rule out the presence of current or past psychiatric illness. Further exclusion criteria for HC were any history of psychiatric illness in first-degree relatives, current or past significant medical or neurological illness. The study was approved by the Ethics Committee of First Affiliated Hospital of Jinan University (China) and the methods and procedures were carried out in accordance with the approved guidelines. All subjects signed a written informed consent form after a full written and verbal explanation of the study. Two senior clinical psychiatrists confirmed that all subjects had the ability to consent to participate in the examination.

MRI data acquisition and preprocessing. MRI data were acquired on a 3.0 T MR system (Discovery MR 750 System, GE Healthcare, Milwaukee, WI) with an 8-channel phased array head coil. Subjects were scanned in a supine, head-first position with symmetrically placed cushions on both sides of the head to decrease motion. During the scanning, participants were instructed to relax with their eyes closed without falling asleep; after the experiment, each participant confirmed not having fallen asleep. The rs-fMRI data were acquired using gradient echo echo-planar imaging sequence with the following parameters: time repetition (TR)/time echo (TE) = 2000/25 ms, slice thickness/gap = 3.0/1.0 mm, voxel size = 3.75 × 3.75 × 3 mm³, field of view (FOV) = 240 × 240 mm, flip angle = 90°, matrix = 64 × 64, number of slices = 35, and NEX = 1. A 7-min scan (210 volumes) was obtained for each subject. In addition, a three dimensional brain volume imaging (3D BRAVO) sequence covering the whole brain was used for structural data acquisition with: TR/TE = 8.2/3.2 ms, slice thickness/gap = 1.0/0 mm, matrix = 256 × 256, FOV = 240 × 240 mm, NEX = 1, flip angle = 12°, bandwidth = 31.25 kHz, and acquisition time = 3 min 45 s. Routine MRI examination images were also collected for excluding anatomic abnormality, such as T1, T2, DWI, and T2- FLAIR images. All participants were confirmed by two experienced radiologists to have no abnormalities on routine MRI.
Functional image preprocessing. The preprocessing was carried out using Data Processing Assistant for Resting-State fMRI (DPARSF)28, which is based on Statistical Parametric Mapping (SPM8, http://www.fil.ion.ucl.ac.uk/spm) and rs-fMRI Data Analysis Toolkit (REST, http://www.restfmri.net)29. For each individual, the first 10 images were discarded to ensure steady-state longitudinal magnetization. The remaining images were slice-time corrected and realigned to the first image in the first series. We controlled for head motion using a threshold of 1.0 mm translation in any cardinal direction and 1.0° rotation in each of the orthogonal x, y, and z axes. The individual T1 images were coregistered to the mean of the realigned EPI images. The coregistered T1 structural images were then segmented (gray matter, white matter, and cerebrospinal fluid) using the unified segmentation algorithm and were then transformed from the individual native space to standard Montreal Neurological Institute (MNI) space. The functional images were also spatially normalized to MNI space by applying the parameters of structural image normalization and were resampled to $3 \times 3 \times 3$ mm$^3$ resolution. The generated images were spatially smoothed with a Gaussian kernel of 4 mm at full width at half-maximum. The resulting fMRI data were linearly trend removed and band-pass (0.01–0.08 Hz) filtered to reduce low-frequency drift and high-frequency noise. Resting-state fMRI measures are sensitive to micro-head motions30, therefore, the Friston 24-Parameter Model30, 31 was used to regress head motion effects from the realigned data (the 24 parameters include six head motion parameters, six head motion parameters one time point before, and the 12 corresponding squared items), based on recent reports demonstrating that higher-order models benefit from the removal of head motion effects30, 31. We further characterized the mean frame-wise displacement (FD), which considers measures of voxel-wise differences in motion in its derivation32, as a measure of the micro-head motion of each subject. Several nuisance variables including mean signals from white matter and cerebrospinal fluid, as well as the mean time series of all voxels across the whole brain, were removed from the preprocessed time-courses by multiple linear regression analysis. Voxel-mirrored homotopic connectivity. VMHC analysis was performed using DPARSF software. A left-right hemisphere symmetric brain template was generated from the all subjects to minimize the effects of geometric differences between the two hemispheres according to Zuo et al’s method17. First, a mean normalized T1 image was generated by averaging all the spatially normalized T1 images. Next, the group-specific symmetrical template was obtained by averaging the mean normalized T1 image and its left-right mirrored version (flipped over x axis). Then each individual T1 image that had been normalized to MNI space was co-registered nonlinearly to this group-specific symmetric template. The same transformation was then applied to the resting-state functional images. For each subject, the homotopic resting state functional connectivity was computed as the Pearson correlation (Fisher Z-transformed) between each voxel’s residual time series and that of its mirrored interhemispheric counterpart. The resultant values generated the VMHC maps and were used for subsequent group-level analyses. We also calculated the global VMHC for each subject and compared it between groups. Global VMHC was calculated by averaging VMHC values across all brain voxels. Statistical analysis. Independent-sample t-tests and chi-squared tests were used to compare demographic data between the RBD and HC groups with SPSS 17.0 software (SPSS, Chicago, IL, USA). All tests were two-tailed, and $P < 0.05$ was considered statistically significant. Nonparametric permutation tests (5,000 permutations) were used to detect statistically significant differences in VMHC between patients with RBD and controls33. Individual’s age, sex, and mean FD were used as nuisance covariates. Multiple comparisons were corrected using threshold-free cluster enhancement (TFCE) ($P < 0.05$)34, which allowed us to avoid arbitrarily choosing a cluster-forming threshold while enabling us to preserve the sensitivity benefits of the cluster-wise correction. Once significant group differences were observed in any brain regions, we further computed Pearson’s correlation coefficients between these VMHC values and the clinical variables including HAMD-24 score, YMRS score, age of illness onset, duration of total illness, duration of symptoms, duration of remissions ($P < 0.05$, Bonferroni corrected), and Spearman’s correlation coefficients between VMHC values and number of episodes ($P < 0.05$, Bonferroni corrected).

Results

Sample characteristics. Table 1 shows the demographic and clinical data of all study participants. Four patients with RBD and four HC were excluded from further analyses due to excessive head motion during the image acquisition. Finally, the participants included 20 patients with RBD and 38 HC subjects. There were no significant differences in sex, age and years of education between the RBD group and the HC group.

VMHC: group differences. The RBD patient group showed lower VMHC values than the control group in the middle frontal gyrus and precentral gyrus (Fig. 1 and Table 2). No region showed greater VMHC in the patient group than in the control group. There were no significant correlations between the VMHC values in these regions and any clinical measures (including HAMD-24 score, YMRS score, age of illness onset, duration of total illness, duration of symptoms, duration of remissions and number of episodes) after bonferroni multiple comparison correction.

Discussion

The VMHC method adopted here to identify brain regions with abnormal connectivity has proved to be informative. Here, VMHC based on the rs-fMRI data was applied for the first time to investigate interhemispheric functional connectivity in patients with RBD. The main finding of this study was the significant decrease in the homologous resting-state functional connectivity in the brain regions of middle frontal gyrus and precentral gyrus in the patients with RBD.

In the current study, we observed decreased VMHC values in the middle frontal gyrus in RBD compared to controls. An rs-fMRI study also found lower VMHC in the medial prefrontal cortex in major depressive disorder35.
The middle frontal gyrus is within the prefrontal lobe and plays an important role in the prefrontal-limbic system. The prefrontal lobe is generally considered to be the advanced cognitive center which participates in integrating emotion and information about internal and external environment, and extracting episodic memory in the resting-state. Two rs-fMRI studies revealed abnormal regional homogeneity in the middle frontal gyrus in BD in the depressive episode. Several researches using task-based fMRI, such as the modified word-based memory task, working memory, response inhibition, and emotional face encoding, indicated decreased or absent activity of the middle frontal gyrus and limbic system in BD patients and increased activity in pediatric BD after pharmacotherapy during a response inhibition task fMRI. Several structural MRI also showed the smaller

| RBD | HC | Statistic |
|-----|----|----------|
| Number of subjects | 20 | 38 | |
| Gender (male/female) | 7/13 | 13/25 | χ² = 0.004, P = 0.952 |
| Age (years) | 28.70 (10.18) | 28.79 (10.18) | t = −0.032, P = 0.975 |
| Education (years) | 13.28 (2.65) | 14.76 (3.13) | t = −1.737, P = 0.09 |
| Age of illness Onset (years) | 21.32 (6.61) | n/a | n |
| Duration of total illness (months) | 59.50 (51.58) | n/a | n |
| Duration of symptoms (months) | 43.64 (48.57) | n/a | n |
| Duration of remissions (months) | 15.86 (18.87) | n/a | n |
| Number of episodes | 3.73 (1.08) | n/a | n |
| HAMD-24 score (points) | 2.59 (1.84) | n/a | n |
| YMRS score (points) | 0.55 (1.30) | n/a | n |
| family history (yes/no) | 4/20 | n/a | n |
| FD values (mm) | 0.09 (0.05) | 0.06 (0.02) | t = 3.687, P = 0.005 |

Table 1. The demographic characteristics of the subjects. Values are reported as mean (standard deviation), except for gender and family history. RBD = remitted bipolar disorder. HC = healthy controls. HAMD-24 = Hamilton Rating Scale for Depression-24. YMRS = Young Mania Rating Scale. FD = framewise displacement for in scanner head motion.

| Brain regions | Brodmann Area | Montreal Neurological Institute Coordinates | Peak Cluster |
|---------------|---------------|-------------------------------------------|--------------|
| Middle frontal gyrus | 6 | X: -33, Y: -6, Z: 57 | t = −3.535, Size: 1026 |
| Precentral gyrus | 4 | X: -36, Y: -18, Z: 51 | t = −4.450, Size: 270 |

Table 2. The areas of VMHC differences between patients with RBD and HC. VMHC = voxel-mirrored homotopic connectivity. RBD = remitted bipolar disorder. HC = healthy controls.
volumes or decreased gray matter in the right middle frontal gyrus in BD. Furthermore, it is interestingly that two of those studies\textsuperscript{46, 45} revealed a consistent result which showed only the volume of the right middle frontal gyrus was inversely correlated with illness duration in BD patients, indicating the middle frontal gyrus may play an important role in the onset of the illness. Studies from literature provide further structural disconnection evidence for functional connectivity abnormalities between the left and right hemispheres of the prefrontal lobe. An international multicenter study showed patients with BD-I had smaller adjusted mid-sagittal CC areas than controls for functional connectivity abnormalities between the left and right hemispheres of the prefrontal lobe. An important role in the onset of the illness. Studies from literature provide further structural disconnective evidence in the left precentral gyrus in BD. Furthermore, numerous structural MRI\textsuperscript{64–66} demonstrated abnormalities in the precentral gyrus in BD, further suggesting the involvement of the precentral gyrus in the pathogenesis of BD. A study\textsuperscript{64} adopted vertex-wise cortical based brain imaging techniques showed BD patients had significantly larger cortical surface area in the left precentral gyrus compared to MDD patients. and a structural MRI\textsuperscript{65} by using Freesurfer technique found increased contrast between grey- and white-matter intensities in the left precentral gyrus in BD, indicating reduced intracortical myelin in this region. A VBM study\textsuperscript{66} found volume deficits in the left precentral gyrus of patients with BD compared to unrelated healthy controls. Intriguingly, van der Schot et al.\textsuperscript{67} also found the genetic risk to develop bipolar disorder was related to decreased grey matter density in the precentral gyrus by investigated 49 affected twin pairs and 67 healthy twin pairs, further suggesting the involvement of the precentral gyrus in the pathogenesis of BD. Therefore, the current study provides further evidence that dysfunction in emotion processing occurs not only in emotional regions of the brain, but also in the traditional motor cortex—precentral gyrus.

The present study showed no significant correlations between the decreased VMHC brain regions and the clinical measures (including HAMD-24 score, YMRS score, age of illness onset, duration of total illness, duration of symptoms, duration of remissions and number of episodes). In a non-clinical depressive symptoms study\textsuperscript{49}, significant correlations between altered ALF values and the depressive severity scores (Beck Depression Inventory scores) was found. In their study, the mean Beck Depression Inventory scores in non-clinical depressive symptoms is 15.24, it met a moderate depression standard according to the Beck Depression Inventory. However, in our study, the mean HAMD-24 score is 2.59 in RBD and this score does not meet any criteria for the depression diagnosis. Thus it’s not hard to understand there is no correlation between the altered VMHC regions and the HAMD-24 score or YMRS score in the RBD patients. It is possible that the alterations of VMHC brain regions (the middle frontal gyrus and precentral gyrus) may be a trait marker\textsuperscript{24} for patients with RBD independent of the severity of symptoms and other related factors, as Guo, W. et al.\textsuperscript{24} reported there is no correlations between the VMHC in any brain region and the overall clinical variables in patients with MDD by voxel-based correlation analyses, and the author speculated that the VMHC abnormal brain regions represented trait characteristics of the disease. Bhagwagar et al.\textsuperscript{69} reported the biochemical abnormalities in recovered unmedicated BD and MDD may be markers of a trait vulnerability to mood disorder, rather than neurochemical correlates of an abnormal mood state. Nevertheless, longitudinal designed study and comparison with the symptomatic BD cohort is necessary to further confirm whether it is a trait marker for BD in future study.

The strengths of the study include the homogeneous samples of RBD. Although the findings of this study are promising, some limitations must be noted. First, the sample size of this study was relatively small, thus, the findings will require replication in a larger study. Second, most of the patients included were under medication prior to MRI scanning. Furthermore, the medication varied, and the number of subjects was too limited to group them according to a specific type of drug. Medication could play a specific role in brain spontaneous activity and functional connectivity but this has not been demonstrated. Studies using patients without medication would be useful for ruling out the effects of medication on the results. Third, the human brain is not symmetrical. However, we tried to solve this issue by applied a symmetrical template and smoothed the functional data to improve the functional correspondence between homotopic regions\textsuperscript{17}. Hence morphometric asymmetry could not account for altered VMHC\textsuperscript{16}. Also, it would be desirable to assess if reduced interhemispheric synchrony observed in patients with RBD is driven by brain activity in a single hemisphere. However, this is not feasible with this functional connectivity approach owing to the operational definition of VMHC analysis, which inhere that results are strictly symmetrical\textsuperscript{17}. Additionally, the underlying mechanisms of altered homotopic connectivity in patients with RBD are largely unknown, it is possible that the structural changes may underlying abnormal interhemispheric synchrony include left-right asymmetries regarding homotopic gray matter volumes. However,
Hermesdorf et al.\textsuperscript{27} investigated the reduced homotopic connectivity in MDD and the underlying structural mechanisms and reported no differences in left-right symmetries in homotopic gray matter volumes were evident across the cohorts. Thus future research that use a functional connectivity combined voxel-based morphometry, Freesurfer and DTI is necessary to clarify the unknown structural basis for VMHC alterations in RBD. Finally, we did not collect neuropsychological data, especially tasks that required interhemispheric transfer of information. Thus, additional neuropsychological tests are needed, and we will investigate the associations between cognitive dysfunction and VMHC deficits in further work.

**Conclusion**

In conclusion, the current findings are the first to reveal aberrant interhemispheric functional connectivity in the bilateral middle frontal gyrus and precentral gyrus in patients with RBD. Since the disruption in functional coordination between the two hemispheres occurred in remission, they may represent trait, rather than state, neurobiological feature of brain function in bipolar disorder. Homotopic interhemispheric resting-state functional connectivity may provide a useful and sensitive screening approach for evaluating BD where neural connectivity is implicated in the pathophysiology.

**References**

1. Geoffroy, P. A. et al. Sleep in remitted bipolar disorder: a naturalistic case-control study using actigraphy. *Journal of affective disorders* **158**, 1–7, doi:10.1016/j.jad.2014.01.012 (2014).

2. Bauer, M. et al. Subsyndromal Mood Symptoms: A Useful Concept for Maintenance Studies of Bipolar Disorder? *Psychopathology* **43**, 1–7, doi:10.1159/000259587 (2010).

3. Torrent, C. et al. Cognitive impairment in bipolar II disorder. *British Journal of Psychiatry* **189**, 254–259, doi:10.1192/bjp.bp.105.071269 (2006).

4. Wang, Y. et al. Interhemispheric resting state functional connectivity abnormalities in unipolar depression and bipolar depression. *Bipolar disorders* **17**, 486–495, doi:10.1111/bdi.12315 (2015).

5. Wang, Y. et al. Reduced interhemispheric resting-state functional connectivity in unmedicated bipolar II disorder. *Acta psychiatraca Scandinavica*. doi:10.1111/acps.12429 (2015).

6. Marlinge, E., Bellivier, F. & Houenou, J. White matter alterations in bipolar disorder: potential for drug discovery and development. *Bipolar disorders* **16**, 97–112, doi:10.1111/bdi.12135 (2014).

7. Sarrazin, S. et al. A Multicenter Tractography Study of Deep White Matter Tracts in Bipolar I Disorder Psychotic Features and Interhemispheric Disconnectivity. *Jama Psychiatry* **71**, 388–396, doi:10.1001/jamapsychiatry.2013.4513 (2014).

8. Wang, F. et al. Abnormal Corpus Callosum Integrity in Bipolar Disorder: A Diffusion Tensor Imaging Study. *Biological psychiatry* **64**, 730–733, doi:10.1016/j.biopsych.2008.06.001 (2008).

9. Emsell, L. et al. Limbic and Callosal White Matter Changes in Euthymic Bipolar I Disorder: An Advanced Diffusion Magnetic Resonance Imaging Tractography Study. *Biological psychiatry* **73**, 194–201, doi:10.1016/j.biopsych.2012.09.023 (2013).

10. Oertel-Knochel, V. et al. Frontal white matter alterations are associated with cognitive function in euthymic bipolar patients. *Journal of affective disorders* **155**, 223–233, doi:10.1016/j.jad.2013.11.004 (2014).

11. Leow, A. et al. Impaired Inter-Hemispheric Integration in Bipolar Disorder Revealed with Brain Network Analyses. *Biological psychiatry* **73**, 183–193, doi:10.1016/j.biopsych.2012.09.014 (2013).

12. Fox, M. D. & Raichle, M. E. Spontaneous fluctuations in brain activity observed with functional magnetic resonance imaging. *Nature Reviews Neuroscience* **8**, 700–711, doi:10.1038/nrn2201 (2007).

13. Vargas, C., Lopez-Jaramillo, C. & Vieta, E. A systematic literature review of resting state network-functional MRI in bipolar disorder. *Journal of affective disorders* **150**, 727–735, doi:10.1016/j.jad.2013.05.083 (2013).

14. Kenny, E. R. et al. Functional Connectivity in Late-Life Depression Using Resting-State Functional Magnetic Resonance Imaging. *American journal of Geriatric Psychiatry* **18**, 643–651, doi:10.1097/GPS.0b013e3181ba2dfe (2010).

15. Salvador, R., Suckling, J., Schwarzbauer, C. & Bullmore, E. T. Undirected graphs of frequency-dependent functional connectivity in whole brain networks. *Philosophical Transactions of the Royal Society B-Biological Sciences* **360**, 937–946, doi:10.1098/rspb.2005.1645 (2005).

16. Kelly, C. et al. Reduced Interhemispheric Resting State Functional Connectivity in Cocaine Addiction. *Biological psychiatry* **69**, 684–692, doi:10.1016/j.biopsych.2010.11.022 (2011).

17. Zuo, X.-N. et al. Growing Together and Growing Apart: Regional and Sex Differences in the Lifespan Developmental Trajectories of Functional Homotopy. *Journal of Neuroscience* **15034–15043, doi:10.1523/neurosci.2612-10.2010 (2010).

18. Zuo, X.-N. & Xing, X.-X. Test-retest reliabilities of resting-state fMRI measurements in human brain functional connectomics: A systems neuroscience perspective. *Neuroscience and Biobehavioral Reviews* **45**, 100–118, doi:10.1016/j.neubiorev.2014.05.009 (2014).

19. Hopman, M. J. et al. Decreased interhemispheric coordination in schizophrenia: A resting state fMRI study. *Schizophrenia research* **141**, 1–7, doi:10.1016/j.schres.2012.07.027 (2012).

20. Li, H.-J., Xu, Y., Zhang, K.-R., Hopman, M. J. & Zuo, X.-N. Homotopic connectivity in drug-naive, first-episode, early-onset schizophrenia. *Journal of Child Psychology and Psychiatry* **56**, 432–443, doi:10.1111/jcpp.12307 (2015).

21. Yuan, K. et al. Reduced Fractional Anisotropy of Corpus Callosum Modulates Inter-Hemispheric Resting State Functional Connectivity in Migraine Patients without Aura. *PloS one* **7**, doi:10.1371/journal.pone.0045476 (2012).

22. Su, Q. J. et al. Decreased interhemispheric functional connectivity in insula and angular gyriusupramarginal gyrus: Significant findings in first-episode, drug-naive somatization disorder. *Psychiatry Research-Neuroimaging* **248**, 48–54, doi:10.1016/j.pscychresns.2016.01.008 (2016).

23. Sun, Y. W. et al. Inter-hemispheric functional and anatomical connectivity abnormalities in traffic accident-induced PTSD: A study combining fMRI and DTI. *Journal of affective disorders* **188**, 80–88, doi:10.1016/j.jad.2015.03.021 (2015).

24. Guo, W. et al. Decreased Interhemispheric Coordination in Treatment-Resistant Depression: A Resting-State fMRI Study. *PloS one* **8**, doi:10.1371/journal.pone.0071368 (2013).

25. Lai, C.-H. & Wu, Y.-T. Decreased inter-hemispheric connectivity in anterior sub-network of default mode network and cerebellum: significant findings in major depressive disorder. *International Journal of Neuropsychopharmacology* **17**, 1935–1942, doi:10.1017/ s1461145714000947 (2014).

26. Wang, L. et al. Interhemispheric Functional Connectivity and Its Relationships with Clinical Characteristics in Major Depressive Disorder: A Resting State fMRI Study. *PloS one* **8**, doi:10.1371/journal.pone.0060919 (2013).

27. Hermosdent, M. et al. Major depressive disorder: Findings of reduced homotopic connectivity and investigation of underlying structural mechanisms. *Human brain mapping* **37**, 1209–1217, doi:10.1002/hbm.23097 (2016).

28. Chao-Gan, Y. & Yu-Feng, Z. DPARSF: A MATLAB Toolbox for “Pipeline” Data Analysis of Resting-State fMRI. *Frontiers in systems neuroscience* **4**, 13–13, doi:10.3389/fnsys.2010.00013 (2010).
29. Song, X. W. et al. REST: A Toolkit for Resting-State Functional Magnetic Resonance Imaging Data Processing. PloS one 6, doi:10.1371/journal.pone.0025031 (2011).

30. Yan, C.-G. et al. A comprehensive assessment of regional variation in the impact of head micromovements on functional connectomics. NeuroImage 76, 183–201, doi:10.1016/j.neuroimage.2013.03.004 (2013).

31. Satterthwaite, T. D. et al. An improved framework for confound regression and filtering for control of motion artifact in the preprocessing of resting-state functional connectivity data. NeuroImage 64, 43–53, doi:10.1016/j.neuroimage.2012.08.052 (2013).

32. Jenkinson, M., Bannister, P., Brady, M. & Smith, S. Improved optimization for the robust and accurate linear registration and motion correction of brain images. NeuroImage 17, 825–841, doi:10.1016/nimg.2002.1132 (2002).

33. Nichols, T. E. & Holmes, A. P. Nonparametric permutation tests for functional neuroimaging: A primer with examples. Human brain mapping 15, 1–25, doi:10.1002/hbm.1058 (2002).

34. Smith, S. M. & Nichols, T. E. Threshold-free cluster enhancement: Addressing problems of smoothing, threshold dependence and localisation in cluster inference. NeuroImage 44, 83–98, doi:10.1016/j.neuroimage.2008.03.061 (2009).

35. Guo, W. et al. Decreased interhemispheric resting-state functional connectivity in first-episode, drug-naive major depressive disorder. Progress in neuro-psychopharmacology & biological psychiatry 41, 24–29, doi:10.1016/j.pnpbp.2012.11.003 (2013).

36. Li, M. et al. Identify Changes of Brain Regional Homogeneity in Bipolar Disorder and Unipolar Depression Using Resting-State fMRI. PloS one 8, doi:10.1371/journal.pone.0079999 (2013).

37. Gao, W. et al. Alterations of regional homogeneity in pediatric bipolar depression: a resting-state fMRI study. BMC Psychiatry 14, doi:10.1186/s12888-014-0222-y (2014).

38. Malhi, G. S. et al. Reduced activation to implicit affect induction in euthymic bipolar patients: An fMRI study. Journal of affective disorders 97, 109–122, doi:10.1016/j.jad.2006.06.005 (2007).

39. Strakowski, S. M., DelBello, M. P. & Adler, C. M. The functional neuroanatomy of bipolar disorder: a review of neuroimaging findings. Molecular Psychiatry 10, 105–116, doi:10.1038/mp.4001585 (2005).

40. Brooks, J. O. III et al. Prefrontal hypooactivation during working memory in bipolar II depression. Psychological medicine 45, 1731–1740, doi:10.1017/s003329171002852 (2015).

41. Penfold, C., Vizueta, N., Townsend, J. D., Bookheimer, S. Y. & Althuler, L. L. Frontal lobe hypooactivation in medication-free adults with bipolar II depression during response inhibition. Psychiatry Research-Neuroimaging 231, 202–209, doi:10.1016/j.pscychresns.2014.11.005 (2015).

42. Tseng, W. L. et al. An fMRI study of emotional face encoding in youth at risk for bipolar disorder. European Psychiatry 30, doi:10.1016/j.eurpsy.2014.05.004 (2015).

43. Pavuluri, M. N., Passarotti, A. M., Harral, E. M. & Sweeney, J. A. Enhanced Prefrontal Function With Pharmacotherapy on a Response Inhibition Task in Adolescent Bipolar Disorder. Journal of Clinical Psychiatry 71, 1526–1534, doi:10.4088/ JCP.q10m05504yel (2010).

44. Frazier, J. A. et al. Cortical grey matter differences identified by structural magnetic resonance imaging in pediatric bipolar disorder. Bipolar disorders 7, 555–569, doi:10.1111/j.1399-5618.2005.00258.x (2005).

45. Ambrosi, E. et al. Structural brain alterations in bipolar disorder II: A combined voxel-based morphometry (VBM) and diffusion tensor imaging (DTI) study. Journal of affective disorders 150, 610–615, doi:10.1016/j.jad.2013.02.023 (2013).

46. Li, M. et al. Voxel-based morphometric analysis on the volume of gray matter in bipolar I disorder. Psychiatry Research-Neuroimaging 191, 92–97, doi:10.1016/j.pscychresns.2010.09.006 (2011).

47. Lyoo, I. K. et al. Regional cerebral cortical thinning in bipolar disorder. Bipolar disorders 8, 65–74, doi:10.1111/j.1399-5618.2006.00284.x (2006).

48. Sarrazin, S. et al. Corpus callosum area in patients with bipolar disorder with and without psychotic features: an international multicentre study. Journal of Psychiatry & Neuroscience 40, 332–339, doi:10.1503/jpn.140262 (2015).

49. Wise, T. et al. Voxel-Based Meta-Analytical Evidence of Structural Disconnection in Major Depression and Bipolar Disorder. Biological psychiatry 79, 293–302, doi:10.1016/j.biopsych.2013.03.004 (2016).

50. Dillon, D. G. & Pizzagalli, D. A. Evidence of successful modulation of brain activation and subjective experience during reappraisal of negative emotion in unmedicated depression. Psychiatry Research-Neuroimaging 212, 99–107, doi:10.1016/j.pscychresns.2013.01.001 (2013).

51. Samson, A. C. et al. Brain activation predicts treatment improvement in patients with major depressive disorder. Journal of psychiatric research 45, 1214–1222, doi:10.1016/j.jpsychires.2011.03.009 (2011).

52. Nitsche, M. A. et al. Facilitation of implicit motor learning by weak transcranial direct current stimulation of the primary motor cortex in the human. J Cogn Neurosci 15, 619–626, doi:10.1162/jocn.2003.15.6.619 (2003).

53. Eksner, C., Koschack, J. & Irle, E. The differential role of premotor frontal cortex and basal ganglia in motor sequence learning: evidence from focal basal ganglia lesions. Learning & memory (Cold Spring Harbor, N.Y.) 9, 376–386, doi:10.1101/lm.48402 (2002).

54. Bunse, T. et al. Motor cortical excitability assessed by transcranial magnetic stimulation in psychiatric disorders: a systematic review. Brain stimulation 7, 158–169, doi:10.1016/j.brs.2013.08.009 (2014).

55. Thomas, L. A. et al. Neural circuitry of masked emotional face processing in youth with bipolar disorder, severe mood dysregulation, and healthy volunteers. Developmental Cognitive Neuroscience 8, 110–120, doi:10.1016/j.dcn.2013.09.007 (2014).

56. Lee, D. et al. Neural correlates of preparatory and regulatory control over positive and negative emotion. Soc Cogn Affect Neurosci 9, 494–504, doi:10.1093/scan/nst115 (2014).

57. Hooker, C. I., Germaine, L. T., Knight, R. T. & D’Esposito, M. Amygdala response to facial expressions reflects emotional learning. Journal of Neuroscience 26, 8915–8922, doi:10.1523/jneurosci.3048-05.2006 (2006).

58. Morata, T. et al. The role of the right prefrontal cortex in self-evaluation of the face: A functional magnetic resonance imaging study. Journal of Cognitive Neurosc 20, 342–355, doi:10.1162/jocn.2008.20024 (2008).

59. Kitada, R., Johnsruade, I. S., Kochiyama, T. & Lederman, S. J. Brain networks involved in haptic and visual identification of facial expressions: An fMRI study. Acta psychiatria Scandinavica 114, 329–338, doi:10.1111/j.1399-5618.2016.12625 (2016).

60. Fung, G. et al. Distinguishing bipolar and major depressive disorders by brain structural morphometry: a pilot study. Bmc Psychiatry 15, doi:10.1186/s12888-015-0685-5 (2015).

61. Jorgensen, K. N. et al. Increased MRI-based cortical grey/white-matter contrast in sensory and motor regions in schizophrenia and bipolar disorders. Psychological medicine 46, 1971–1985, doi:10.1017/s0033291716000593 (2016).

62. Eker, C. et al. Brain regions associated with risk and resistance for bipolar I disorder: a voxel-based MRI study of patients with bipolar disorder and their healthy siblings. Bipolar disorders 16, 249–261, doi:10.1111/bdi.12181 (2014).

63. van der Schot, A. C. et al. Genetic and environmental influences on focal brain density in bipolar disorder. Brain: a journal of neuroscience 133, 3080–3092, doi:10.1093/brain/awq236 (2010).
68. Wei, X. et al. Alteration of spontaneous neuronal activity in young adults with non-clinical depressive symptoms. Psychiatry research 233, 36–42, doi:10.1016/j.psychres.2015.04.008 (2015).

69 Bhagwagar, Z. et al. Reduction in Occipital Cortex GABA Concentrations in Medication-Free Recovered Unipolar Depressed and Bipolar Subjects. Biological Psychiatry 61(6), 806–812 (2007).

Acknowledgements
We would like to thank all the participants for their contribution to this study. The study was supported by grants from the National Natural Science Foundation of China (81671670, 81501456, 81471650); Planned Science and Technology Project of Guangdong Province, China (2014B020212022); Planned Science and Technology Project of Guangzhou, China (1563000653, 201508020004, 20160402007, 201604020184). The funders had no role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Author Contributions
Ying Wang, Lianping Zhao and Li Huang were involved in the design of this study and contributed to the writing of the manuscript. Zhangzhang Qi performed the MR scan. Lianping Zhao, Ying Wang and Yao Sun performed the MR data analyses. Yanbin Jia and Shuming Zhong were involved in diagnosing subjects and clinical data collection. Lianping Zhao and Ying Wang wrote the first draft of the manuscript. Zhongping Zhang provided the revisions of the data processing. Li Huang provided critical revisions of the manuscript. All authors approved the manuscript for submission. The authors of this paper do not have any commercial associations that might pose a conflict of interest in connection with this manuscript.

Additional Information
Competing Interests: The authors declare that they have no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2017