Analysis of Altered Baseline Brain Activity in Drug-Naive Adult Patients with Social Anxiety Disorder Using Resting-State Functional MRI

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Objective We hypothesize that the amplitude of low-frequency fluctuations (ALFF) is involved in the altered regional baseline brain function in social anxiety disorder (SAD). The aim of the study was to analyze the altered baseline brain activity in drug-naive adult patients with SAD.

Methods We investigated spontaneous and baseline brain activities by obtaining the resting-state functional magnetic resonance imaging data of 20 drug-naïve adult SAD patients and 19 healthy controls. Voxels were used to analyze the ALFF values using one- and two-sample t-tests. A post-hoc correlation of clinical symptoms was also performed.

Results Our findings show decreased ALFF in the bilateral insula, left medial superior frontal gyrus, left precuneus, left middle temporal gyrus, right middle temporal pole, and left fusiform gyrus of the SAD group. The SAD patients exhibited significantly increased ALFF in the right inferior temporal gyrus, right middle temporal gyrus, bilateral middle occipital gyrus, orbital superior frontal gyrus, right fusiform gyrus, right medial superior frontal gyrus, and left parahippocampal gyrus. Moreover, the Liebowitz Social Anxiety Scale results for the SAD patients were positively correlated with the mean Z values of the right middle occipital and right inferior occipital but showed a negative correlation with the mean Z values of the right superior temporal gyrus and right medial superior frontal gyrus.

Conclusion These results of the altered regional baseline brain function in SAD suggest that the regions with abnormal spontaneous activities are involved in the underlying pathophysiology of SAD patients.

Key Words Social anxiety disorder, Amplitude of low-frequency fluctuations, Default mode network.

INTRODUCTION

Social anxiety disorder (SAD), also known as social phobia, is the persistent fear of social or performance situations in which the person is exposed to unfamiliar people or to possible public scrutiny.1 Resting-state functional magnetic resonance imaging (fMRI) is a novel technique that has several potential advantages over task performance or stimulation in terms of clinical applicability.2-4 Previous studies have found altered brain functions in the medial prefrontal cortex (MPFC) and limbic regions, including the amygdala, hippocampus, and insula, of SAD patients.5-13 However, investigation on SAD at the baseline levels or studies on the spontaneous activity of the brain must be conducted to obtain a clearer understanding of the neurobiology of the disorder.1,14 Furthermore, a relationship has been observed between the altered spontaneous activity of the functionally relevant cortices and clinical features.14 Thus, studying the spontaneous cerebral
activity in drug-naïve adult patients with SAD may provide vital information for characterizing the neuropathophysiological mechanisms underlying the disorder.

The amplitude of low-frequency fluctuations (ALFF), which is a widely accepted resting-state data analysis tool that can provide information on regional spontaneous neuronal activity (SNA), has been extensively used to study mental disorders. Brain diseases may exhibit abnormal local SNA and/or inter-regional SNA synchronization while ALFF reflects the extent of SNA. Hence, the changed SNA of these regions by ALFF may be implicated in the underlying pathophysiology in disease. Thus, this method has been suggest to investigate the functional modulations and characterize the neuropsychological changes in the resting state in patients with various clinical populations. Although several studies have demonstrated the altered resting-state functional connectivity in various brain networks during SAD, to the best of our knowledge, no resting-state fMRI study has been conducted on the ALFF in SAD. By using resting-state fMRI and ALFF analysis, the current research focused on the local coherence of spontaneous activity of drug-naïve patients with SAD. This study aims to determine whether 1) the ALFF in some brain areas of SAD patients are aberrant, and 2) these changes are related to the measured clinical severity of the disorder.

METHODS

Subjects
Twenty patients (22.90±2.99 years old, all right-handed) were consecutively recruited from West China Hospital of Sichuan University. A diagnosis of SAD was determined by consensus between two attending psychiatrists using the Structured Clinical Interview for DSM-IV (SCID) Patient Edition. The exclusion criteria included history of neurological and psychiatric disease and diagnosis of other mental disorders, except SAD; existence of an organic brain disorder; alcohol or drug abuse; and pregnancy or any physical illness such as hepatitis, brain tumor, or epilepsy, as assessed from the medical records. No gross abnormalities were observed in the brain MRI scans (i.e., T1-weighted and T2-weighted images) of any of the subjects when examined by an experienced neuroradiologist. The SAD patients have yet to receive psychotherapy and psychiatric medications.

Twenty age- and education-matched healthy controls (HCs) (mean age=21.38±3.77 years, all right-handed) were recruited from the local community via poster advertisement and then screened using the SCID-I/P version to confirm the absence of any psychiatric or neurological illness. The HC subjects were told beforehand to cooperate in the radiological check for fear of intentional withdrawal because of some mental disorders. All HC subjects were interviewed to confirm that no history of psychiatric illness exists among their first-degree relatives. However, one person was involuntarily discontinued during MR scanning because of hypertension. Finally, image data were acquired from 19 HCs.

The participants of the two groups were evaluated using the Liebowitz Social Anxiety Scale (LSAS), State-Trait Anxiety Inventory (STAI), the Hamilton Anxiety Rating Scale (HAMA), and Hamilton Depression Rating Scale (HAMD). According to previous studies, LSAS does not replace a clinical interview for the diagnosis of social anxiety even when the score exceeds 38, which is indicative a probable SAD diagnosis. The present study was approved by the local Ethics Committee. A written informed consent was obtained from all subjects after they were provided with a complete description of the study.

Image acquisition
Experiments were performed using a 3.0-T GE-Sigma MRI scanner (EXCITE, General Electric, Milwaukee, WI, USA). A foam padding was used to minimize head motion. Functional images were acquired using a single-shot, gradient-recalled echo-planar imaging sequence (TR=2000 ms, TE=30 ms, and flip angle=90°). Thirty transverse slices (FOV=24 cm, in-plane matrix=64×64, slice thickness=5 mm, without gap) aligned along the anterior commissure-posterior commissure line were acquired. A total of 205 volumes were acquired for each subject, and the first 5 volumes were discarded to ensure steady-state longitudinal magnetization. Subsequently, for spatial normalization and localization, a set of high-resolution T1-weighted anatomical images in the axial orientation were acquired for each subject using a 3D spoiled-gradient recalled sequence (TR=8.5 ms, TE=3.4 ms, flip angle=12°, matrix size=512×512×156, and voxel size=0.47×0.47×1 mm³).

Data preprocessing
Data preprocessing was partly conducted using the Statistical Parametric Mapping software (SPM2, http://www.filion.ucl.ac.uk/spm). The 200 volumes were first corrected for temporal difference in the acquisition of the different slices. Afterward, the images were realigned to the first volume for head-motion correction. For one subject, the translational or rotational parameters in these datasets exceeded ±1.5 mm or ±1.5°; these datasets were thus excluded from the analysis. We also evaluated the group differences in translation and rotation of head motion according to the following formula: Head Motion/Rotation = \frac{1}{L-1} \sum_{i=1}^{L} \sqrt{(x_i-x_{i-1})^2+(y_i-y_{i-1})^2+(z_i-z_{i-1})^2},

where L is the length of the time series (L=200 in this study), x, y, and z are translations/rotations at the ith time point in the x, y and z directions, respectively. The fMRI images were
realigned with the corresponding T1 volume and warped into a standard stereotaxic space at a resolution of $3\times3\times3$ mm$^3$ using the Montreal Neurological Institute echo-planar imaging template in SPM2. Afterward, the images were spatially smoothed via convolution with an isotropic Gaussian kernel (FWHM=8 mm).

**ALFF analysis**

The ALFF analysis was performed using the REST software (http://resting-fmri.sourceforge.net). The following procedure was applied to calculate the ALFF: after band pass filtering (0.01 Hz to 0.08 Hz) and linear-trend removal, the time series was transformed into a frequency domain using fast Fourier transform (FFT) (taper percent: 0, FFT length: shortest), and the power spectrum was obtained. Given that the power of a given frequency is proportional to the square of the amplitude of this frequency component in the original time series in the time domain, the square root of the power spectrum obtained by FFT was calculated and then averaged across 0.01 Hz to 0.08 Hz at each voxel. This averaged square root was taken as the ALFF. For normalization purposes, the ALFF of each voxel was divided by the mean ALFF value within a brain mask.

**Statistics and post-hoc correlation analysis**

A one-sample, one-sided t-test was performed within each group to determine whether the ALFF deviated from the value of 1. A two-sample t-test was performed to verify the ALFF difference between the two groups. Voxels ($p<0.001$, uncorrected; cluster size >10 voxels) were used to show the significant difference between the two groups. Two-sample t-tests were also performed to assess the differences in the head motions of the two groups. A post-hoc Pearson correlation analysis was also performed to investigate the relationship between the ALFF values and clinical severity. Given that these analyses were exploratory in nature, a statistical significance level of $p<0.05$ (uncorrected) was used.

**RESULTS**

**Neuropsychological data and group characteristics**

The group demographic characteristics and neuropsychological scores are shown in Table 1. No significant difference was found in the gender, age, and education of the different groups. The LSAS total score ($t=11.02$, $p<0.001$), fear factor ($t=10.90$, $p<0.001$), and avoidance factor ($t=7.88$, $p<0.001$) of the SAD patients were significantly higher than those of the healthy controls (HCs). Significant differences were also

### Table 1. Neuropsychological data and group characteristics

|                          | SAD (N=20) | HC (N=19) | SAD vs. HC |
|--------------------------|-----------|-----------|------------|
|                          | M±SD      | M±SD      | t value    | p value    |
| Gender (n: male/female)  | 14M/6F    | 14M/5F    | -          | -          |
| Age (years)              | 22.90±3.99| 21.79±3.61| 0.91       | 0.369      |
| Education (years)        | 14.10±1.48| 14.11±2.00| -0.01      | 0.993      |
| Duration (months)        | 45.40±39.78| -         | -          | -          |
| LSAS                     |           |           |            |            |
| Total score              | 53.90±11.50| 19.21±7.68| 11.02      | <0.0001    |
| Fear factor              | 28.00±6.17 | 8.42±4.94 | 10.90      | <0.0001    |
| Avoidance factor         | 25.90±6.93| 10.79±4.79| 7.88       | <0.0001    |
| HAMD                     | 7.50±6.27  | 1.05±1.54 | 4.36       | <0.001     |
| HAMA                     | 7.50±6.27  | 0.89±1.52 | 4.65       | <0.0001    |
| STAI                     |           |           |            |            |
| STAI-T                   | 48.25±7.02 | 32.58±4.85| 8.07       | <0.0001    |
| STAI-S                   |           |           |            |            |
| Pre-scanning             | 41.35±8.31 | 31.05±4.72| 4.73       | <0.0001    |
| Post-scanning            | 37.65±9.54 | 32.68±7.02| 1.84       | 0.073      |
| Head Motion              |           |           |            |            |
| Translation (mm)         | 0.03      | 0.01      | 0.04       | 0.02       |
| Rotation (°)             | 0.03      | 0.01      | 0.03       | 0.02       |

Data from questionnaires are presented in terms of mean score (M) and standard deviation (SD) in SAD and HC groups. The p values were obtained by two-sample two-tailed t-test. SAD: social anxiety disorder, HC: healthy control, LSAS: Liebowitz Social Anxiety Scale, HAMA: Hamilton Anxiety Rating Scale, HAMD: Hamilton Depression Rating Scale, STAI: State-Trait Anxiety Inventory, T: trait, S: state.
found between HAMD (t=4.36, p<0.001) and HAMA (t=4.65, p<0.001) of the two groups. Compared with the HCs, the SAD patients showed significantly higher levels of anxiety, as assessed by STAI-T (t=8.07, p<0.001) and prescanning STAI-S (t=4.73, p<0.001). No difference was found in the post-scanning STAI-S (t=1.84, p=0.073) of the patients and the HCs. Moreover, no significant head motion difference was observed between the two groups (shift: t=-1.15, p=0.259; rotation: t=-0.95, p=0.351) (Table 1).

Activity pattern in each group and altered ALFF in SAD patients

The one-sample t-tests of the ALFF analysis showed significantly higher ALFF in the posterior cingulate cortex/precuneus (PCC/Pcun), MPFC, and bilateral angular gyrus of the HCs and SAD patients (p<0.001, uncorrected) (Figure 1). The two-sample t-test showed significant difference in some brain areas of the two groups (p<0.001, uncorrected) (Figure 2). In the SAD group, the brain regions with decreased ALFF included the bilateral rolandic operculum, postcentral gyrus, superior temporal gyrus, middle frontal gyrus, triangular inferior frontal gyrus, supramarginal gyrus, superior parietal gyrus, and insula (Table 2, Figure 2). Conversely, the SAD patients exhibited significantly increased ALFF in these cortices, including the right inferior temporal gyrus and right middle temporal gyrus (these two areas were merged into one cluster), as well as in the bilateral middle occipital gyrus, orbital superior frontal gyrus, right inferior occipital gyrus, right orbital medial frontal gyrus, and other re-
Correlation of the regions of decreased ALFF with the anxiety scale of patients

The LSAS values for the SAD patients showed a significantly positive correlation with the mean Z values of the right middle occipital and right inferior occipital but was negatively correlated with those of the right superior temporal gyrus and right medial superior frontal gyrus (p<0.05) (Figure 3). In addition, the other brain regions with aberrant ALFF showed no significant correlation with the LSAS values.

DISCUSSION

To the best of our knowledge, the present study is the first to report on an association between ALFF at rest and the severity of social anxiety symptoms in patients, particularly drug-naive adults. We found a significant difference in the ALFF of the SAD patients and the HCs. The SAD group showed altered ALFF in insula, medial superior frontal gyrus, PCC/Pcun, temporal lobe and fusiform gyrus, and occipital lobe. In addition, the LSAS values of SAD patients showed a positive correlation with the mean Z values of the right middle occipital and right inferior occipital but negatively with that of the right superior temporal gyrus and right medial superior frontal gyrus.

Changes in the anxiety level during post-scanning

Both STAI-T and prescanning STAI-S results showed that the SAD patients had higher anxiety levels than HCs, whereas

Table 2. Regions showing decreased ALFF in the subjects between SAD and HC groups

| Anatomical region             | Hemisphere | MNI coordinates (x, y, z) | Brodmann’s area | Cluster size (voxels) | t value |
|-------------------------------|------------|--------------------------|-----------------|-----------------------|---------|
| SAD<HC                        |            |                          |                 |                       |         |
| Rolandic operculum            | L          | -48 -18 12              | 6,22,48         | 153                   | -4.19   |
|                               | R          | 66 -12 12               | 22,48           | 63                    | -4.82   |
| Postcentral gyrus             | L          | -63 -15 18             | 22,42,43        | 94                    | -4.74   |
|                               | R          | 69 -15 15              | 1,72,22         | 56                    | -5.05   |
| Superior temporal gyrus       | L          | -51 -18 12             | 22,42,48        | 70                    | -4.83   |
|                               | R          | 69 -15 12              | 22,41,48        | 64                    | -4.71   |
| Superior frontal gyrus, medial| L          | -9 48 39              | 8,9,32          | 68                    | -3.75   |
| Inferior parietal gyrus       | L          | -51 -54 45             | 7,39,40         | 64                    | -3.72   |
| Insula                        | L          | -45 0 6               | 48              | 60                    | -3.80   |
|                               | R          | 39 -15 12             | 48              | 40                    | -3.45   |
| Inferior frontal gyrus, orbital| R          | 45 39 -6              | 11,47           | 48                    | -4.35   |
| Precuneus                     | L          | -6 -45 48              | 5               | 45                    | -4.33   |
| Middle frontal gyrus          | L          | -24 48 33             | 44,46           | 10                    | -3.85   |
|                               | R          | 39 27 21              | 9,45,46         | 43                    | -3.29   |
| Superior frontal gyrus        | R          | 21 -3 75              | 6,8,9           | 41                    | -3.5776 |
| Inferior frontal gyrus, triangular| L          | -54 18 0              | 45              | 18                    | -3.37   |
|                               | R          | 45 39 -3              | 45,47           | 40                    | -3.87   |
| Middle temporal gyrus         | L          | -66 -27 0             | 20,21,22        | 34                    | -3.09   |
| Heschl gyrus                  | L          | -33 -24 6             | 22,48           | 31                    | -4.17   |
| Middle temporal gyrus, pole   | R          | 51 6 -24             | 20,21,28        | 21                    | -3.39   |
| Precentral gyrus              | R          | 57 0 21              | 6               | 19                    | -3.38   |
| Median cingulate gyrus        | L/R        | -6 -45 48             | 23              | 25                    | -4.33   |
| Supramarginal gyrus           | L          | -54 -21 15            | 22,42,48        | 14                    | -3.70   |
|                               | R          | 69 -18 18             | 20,40,48        | 18                    | -3.75   |
| Superior parietal gyrus       | L          | -33 -72 51            | 7               | 10                    | -3.35   |
|                               | R          | 24 -45 78             | 5,7             | 10                    | -3.22   |
| Fusiform gyrus                | L          | -33 -39 -15           | 37              | 4                     | -2.95   |

The threshold was at p<0.001, uncorrected by two-sample t-test. ALFF: amplitude of low-frequency fluctuations, SAD: social anxiety disorder, HC: healthy control, MNI: Montreal Neurological Institute, L: left, R: right
post-scanning STAI-S showed no significant differences between the anxiety levels of the two groups. This finding is consistent with those of previous studies. SAD patients often abnormally feel anxious and worried about embarrassing themselves in unfamiliar situations. This mental state is different from scan-related anxiety. The clinical scale results suggest that the symptoms of SAD patients change with the social situation; that is, these patients do not develop anxiety symptoms until they find themselves in a social setting. The decrease in the post-scanning anxiety level indicates that anxiety anticipation reached remission and was released when the subjects were removed from the social setting. Thus, anxi-

Table 3. Regions showing increased ALFF in the subjects between SAD and HC groups

| Anatomical region                  | Hemisphere | MNI coordinates (x, y, z) | Brodmann’s area | Cluster size (voxels) | t value |
|-----------------------------------|------------|---------------------------|-----------------|-----------------------|---------|
| Superior temporal gyrus           | R          | 42 -42 -12                | 20,37           | 47                    | 3.57    |
| Middle temporal gyrus             | R          | 54 -57 0                 | 37              | 44                    | 3.12    |
| Middle occipital gyrus            | L          | -36 -87 18               | 18,19           | 40                    | 3.54    |
|                                   | R          | 42 -87 0                 | 18,19           | 39                    | 3.53    |
| Superior frontal gyrus, orbital   | L          | -21 48 -15               | 11              | 20                    | 3.44    |
|                                   | R          | 18 48 -15                | 11              | 29                    | 3.42    |
| Inferior occipital gyrus          | R          | 42 -87 -3                | 18,19           | 22                    | 3.34    |
| medial frontal gyrus, orbital     | R          | 15 -42 -3               | 10,11           | 22                    | 3.13    |
| Fusiform gyrus                    | R          | 42 -45 -12              | 20,37           | 4                     | 3.09    |
| Superior frontal gyrus, medial    | R          | 18 -45 0                 | 10,11           | 4                     | 2.89    |
| Parahippocampal gyrus             | L          | -24 -42 -3              | 37              | 3                     | 2.84    |

The threshold was at p<0.001, uncorrected by two-sample t-test. ALFF: amplitude of low-frequency fluctuations, SAD: social anxiety disorder, HC: healthy control, MNI: Montreal Neurological Institute, L: left, R: right.

Figure 3. Results of the correlation analysis between LSAS (LSAS-F: fear factor, LSAS-A: avoidance factor) values and ALFF in the SAD group. This scatter diagram shows the positive and negative correlations of LSAS with the mean Z values of the altered ALFF regions in SAD patients at the threshold of p<0.05 via post-hoc correlation analysis. LSAS: Liebowitz social anxiety scale, ALFF: amplitude of low-frequency fluctuations, SAD: social anxiety disorder, MNI: Montreal Neurological Institute.
ety anticipation is vital to the pathopsychological mechanism of SAD. However, this finding requires further verification by a combined study of tasking-state fMRI and behavioral tests.

Insula
Our results show decreased ALFF in the insula of SAD patients. By contrast, a number of studies reported that hyperfunction occurs more frequently than hypofunction in insula. In accordance with previous studies, we hypothesize that the persistent hyperactivity in the insula of anxious patients disrupts the spontaneous neural activity and function for anticipatory processing in the insula. These differences in insula activity, which range from tasking to the resting state, may be explained by the implicit sequence learning during the resting state; this type of learning is a cognitive task that is not aimed at symptom provocation. In addition, this area may be critical for the extraction and selection of task-relevant information and has been implicated in inhibitory control. Thus, the insula showed hyperfunction during symptom provocation. Moreover, the capacity of this brain region to deal with a cognitive task even in the resting state was disrupted.

PCC/Pcun and default mode network
The one-sample t-tests in the ALFF analysis showed significantly higher ALFF in the PCC/Pcun, MPFC, and bilateral AG of the HCIs (Figure 1A). This pattern is consistent with that of the DMN. In social cognition and "theory of mind", the PCC/Pcun is sometimes referred to as a pivotal hub of the DMN. Moreover, the PCC/Pcun is more active when socially relevant emotional stimuli are perceived and is involved in a network of self-consciousness and self-related mental representations. Social cognition and "theory of mind", which are two preeminent cognitive behavioral models of social anxiety, have described that self-focused attention in SAD and misinterpretation of interoceptive information increase access to negative thoughts and feelings; these negative reactions prevent individuals from gaining accurate information (about the situation and the responses of others) that could diffuse the negative expectations. Thus, our study provided further evidence that in SAD, Pcun dysfunction results in inaccurate evaluation of the self-emotional state as well as in incomplete processing of self-related representations. Given the role of PCC/Pcun in state perception and attribution, our findings support the hypothesis that PCC/Pcun can suspend its function within the DMN. We propose that the impairment of PCC/Pcun in the DMN of SAD patients may be relevant in the development of the feeling of wariness of others' judgment as well as in the remaining high-level, self-focused attention.

Frontal lobe
The aberrant ALFF of SAD patients are represented in the right medial superior frontal gyrus. In previous studies, the activities in the MPFC of SAD subjects either increased or decreased but showed no significant difference during task performance. The MPFC activity possibly reflects an interaction between cognitive processing and emotional state. Meanwhile, the prefrontal cortex activation appears to correlate with the self-referential magnitude and higher evaluation processes, particularly with the construction of self-relevant mental simulation in the form of memories. Our findings, along with the hypotheses on MPFC, suggest that MPFC is significantly involved in the prominent psychopathological mechanism of SAD. Interestingly, the orbitofrontal/MPFC has been hypothesized to have a role in the inhibition or extinction of excessive corticolimbic activity in anxiety disorders. However, our results show no evidence of excessive amygdala-hippocampal activity in the patients during the resting state. The more likely frontal compensation, does not prevent limbic overactivity, as shown by the aberrant ALFF in the resting-state frontal cortex. Whether this result indicates a compensatory mechanism in the brain requires further research. Meanwhile, the prefrontal cortex may be involved in the internal representation of the social environment and may also have a role in our ability to read the mental states of other people. The frontal dysfunction from this study possibly reflect the dysfunction of the cortical regions that directly contribute to the etiology of SAD. Our results on the bilateral medial superior frontal gyrus have not been previously reported. These results are associated with the relationship between the SAD subtype and other factors and thus require further investigation.

Fusiform and temporal lobe
In this study, decreased ALFF was observed in the temporal cortex, including the bilateral superior temporal gyrus, superior parietal gyrus, left inferior parietal gyrus, left middle temporal gyrus, and right middle temporal pole. These results are inconsistent with the significantly increased ALFF in the right inferior/middle temporal gyrus and right fusiform gyrus (Figure 2, Table 2) but partly agree with the results of previous tasking-state studies on the fusiform of SAD subjects. The fusiform can form a higher-sensory cortex, which is involved in the perceptual representation of social stimuli. Alternatively, the increased ALFF in the fusiform gyrus may be due to the higher vigilance and alertness of SAD patients, whereas the decreased ALFF in the fusiform gyrus may be due to avoidance. This anterior interpretation is supported by the relationships between the mean Z values of ALFF in the right superior temporal gyrus, whereas the LSAS results were
mostly correlated with the fear factor.\textsuperscript{35}

The negative correlation between the mean $Z$ values of ALFF in the right superior temporal gyrus and the LSAS scores further provides evidence for the functional impairments of SAD patients in terms of resting-state facial perception and social evaluation. Although the function of the temporal poles is not well understood, damaged temporal poles can impair the ability to use the knowledge we gain from the world through our experiences; this knowledge is important for our ability to mentalize.\textsuperscript{44} Thus, the abnormal ALFF values in the aforementioned brain regions indicate a functional defect in the SAD patients to evaluate the current situation correctly using their experience and/or by observing the behavior of others. The functional imaging results suggest that temporal cortex dysfunction has a vital function in the psychopathology of SAD.

Occipital lobe

In this study, the significantly reduced ALFF in the cortices, including the pre- and post-central gyrus, and increased ALFF in the middle and inferior occipital gyriuses contribute to perceptual impairments of the sensorimotor cortex (SMC) and supersensitivity of the visual cortex of SAD patients. Some of these results are consistent with the findings of previous tasking-state fMRI studies and suggest that sensorimotor and the visual cortex showed significantly stronger BOLD responses to social threat in SAD patients.\textsuperscript{7} Moreover, the significant positive correlation between the LSAS scores and the mean ALFF $Z$ values in the right middle occipital gyrus and right inferior occipital gyrus of the patients further provide evidence for the resting-state hyperactivity of the visual cortex in SAD. Along with our current data, these functional imaging results suggest a dysfunction in the SMC area and visual cortex of SAD patients. Thus, the hypothesis that SAD patients are over-aroused and exhibit hyperprosexia for social information\textsuperscript{41} as a result of hyperactivity in the visual cortex was confirmed.

Several limitations must be considered in this research. First, the amygdala has a vital function in fear. However, in this study, we did not find significant differences in the ALFF of the amygdala of the two groups, possibly because of the relatively low fMRI signal-to-noise ratio in the basal ganglia in resting-state fMRI. Second, head motion is relatively a greater issue for anxiety-disorder patients than for the controls during fMRI. In this study, we did not find significant differences in the head motion of the two groups. However, a more restricted head motion should yield more accurate and more reliable results. Third, the physiological noise induced by cardiac and respiratory rhythms is a significant factor in functional neuroimaging studies, particularly in persons with anxiety disorders.\textsuperscript{15} In this study, we were unable to resolve the bias generated by these noises. Future studies could focus on a few specific regions of interest and utilize multiple research methods, such as resting- and tasking-state fMRI and PET. In addition, a larger group of subjects would increase the statistical power of the study.

Conclusions

Our findings indicate differential impairments or dysfunctions in the neural activity of the insula, PCC/Pcun, and frontal and temporal cortices in SAD patients. These results suggest that the regions with abnormal spontaneous activities are involved in the underlying pathophysiology of SAD patients. These findings also support the results of a tasking-state study, which showed that abnormal responses may be partly due to abnormal spontaneous baseline neural activity.

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