Associations of cortical thickness, surface area and subcortical volumes with insight in drug-naïve adults with obsessive-compulsive disorder

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ABSTRACT

Poor insight in obsessive-compulsive disorder (OCD) is associated with several adverse clinical outcomes. However, the neurobiological basis of this insight deficit is not clearly understood. The present study thus aimed to investigate associations of cortical thickness, cortical surface area and subcortical volumes with insight in a sample of drug-naïve adults with OCD. Forty-seven OCD patients and 42 healthy controls (HCs) underwent MRI scanning. Cortical thickness and surface area were compared on a vertex-by-vertex basis across groups, while subcortical volumes were compared on a structure-by-structure basis. Partial correlation analyses were then performed to assess associations between regional cortical and subcortical measures and insight levels. OCD-GI and OCD-PI groups displayed partly shared, but also partly distinct brain structural alterations. Strikingly, OCD-PI showed decreased cortical thickness in the left superior frontal gyrus, left anterior cingulate cortex (ACC) and right inferior parietal gyrus, compared to both OCD-GI and HCs. Average cortical thickness extracted from these areas was further negatively correlated with BABS scores in the OCD-PI patients. Our findings suggest that poor insight in patients with OCD may have a neural substrate involving the left medial frontal and the right inferior parietal cortices.

1. Introduction

It is generally accepted that poor insight is a hallmark of psychotic disorders (Amador and David, 2004), the clinical significance of which was underscored due to its close relationship with treatment non-adherence (Leclerc et al., 2015). For many years, the study of insight has been primarily centered on schizophrenia (Mintz et al., 2003; Pousa et al., 2017), and only very recently it has become a focus of interest in obsessive-compulsive disorder (OCD). Up to 15–36% of OCD patients demonstrated poor insight into their symptoms and pathology (Catapano et al., 2010; Matsunaga et al., 2002). Importantly, poor insight in OCD has been linked with an array of less favorable clinical outcomes, including more severe OC symptoms (Bellino et al., 2005; Catapano et al., 2001; Ravi et al., 2004), lower age at illness onset (Catapano et al., 2010; Matsunaga et al., 2002., Ravi et al., 2004), higher rates of psychiatric co-morbidity with depression (Ravi et al., 2004) and body dysmorphic disorder (Eisen et al., 2004), and insufficient responses to both behavior therapy (Himle et al., 2006; Mataix-Cols et al., 2002) and pharmacological interventions (Catapano et al., 2001, 2010; Erzagovesi et al., 2001; Ravi et al., 2004), all of which are believed to contribute to a worse prognosis. These results promote research efforts to elucidate the neurobiological mechanisms that underlie poor insight in OCD.

Several studies suggested that insight in OCD has a structural brain basis. Evidence has come from studies examining neuropsychological correlates of poor insight in OCD and then detecting a significant relationship between poor insight and worse cognitive performance, such as executive function and verbal memory (Kashyap et al., 2012;
Kitis et al., 2007; Tumkaya et al., 2009), which are in turn associated with volume of specific brain regions (Choi et al., 2004; Christian et al., 2008). Compared to their good insight counterparts, OCD patients with poor insight demonstrated more severe neurological soft signs, indicating more extensive neurodevelopmental abnormalities in this subgroup (Karadag et al., 2011), which could be reflected in changes of brain morphology. Aigner et al. (2005) directly compared Magnetic Resonance Imaging (MRI) findings between OCD patients with good and poor insight and found that the poor insight cohort showed a higher frequency of structural abnormalities. However, imaging findings in this study were qualitatively evaluated as abnormal or normal, from which we can only infer that poor insight would be at the more “organic” end of the spectrum. The neuroanatomical basis of poor insight in OCD remains largely unknown.

In recent years, surface-based morphometry (SBM) methods have been increasingly applied to investigate the neuroanatomical mechanisms of psychiatric disorders. Unlike the traditional voxel-based morphometry (VBM), which involves a generic voxel-wise comparison of regional concentration or volume of gray matter between different groups (Ashburner and Friston, 2000), SBM utilizes cortical geometric models to partition its constituent cortical thickness and surface area components, which are thought to have differing genetic determinants, phylogeny, and development trajectories (Armstrong et al., 1995; Joyner et al., 2009; Panizzon et al., 2009; Wierenga et al., 2014). It has been proposed that measures of cortical thickness and surface area reflect distinct aspects of the underlying neural architecture. For instance, according to the radial unit hypothesis, the cerebral cortex is organized into ontogenetic columns, and cells within a column have a common origin before moving to their location within the cortex during development (Rakic et al., 1988). In this model, cortical thickness is a marker for the number of cells within columns, whereas surface area primarily reflects the number of cortical columns (Rakic, 1988, 1995, 2007). It is easy to understand that reduction of cells in columns could first result in a thinner thickness but don’t have to cause a significant decrease in surface area or volume. Therefore, surface-based analysis, especially the cortical thickness, has been proposed to be a more useful measure to assess the neuroanatomical patterns associated with the neurodegenerative process and a more biologically informative approach with particular sensitivity in capturing subtle, previously undetected anatomical changes (Kelly et al., 2013; Pereira et al., 2012).

In previous SBM studies, cortical abnormalities were well documented to be involved in the pathophysiology of OCD and have been linked to OCD-specific clinical characteristics (Fan et al., 2013; Fouche et al., 2017; Fullana et al., 2014; Kühn et al., 2013; Nakamae et al., 2012; Shin et al., 2007; Venkatasubramanian et al., 2012). A recent meta- and mega-analysis argued that cortical thickness and surface area represent distinct morphological features and may be differentially affected by different stages of illness and disease profiles of OCD (Boedhoe et al., 2017). Notably, emerging evidence has linked thickness reduction in the frontal, parietal as well as temporal regions with poor insight in schizophrenia and psychosis (Buchy et al., 2012, 2017; Emami et al., 2016). Though not examined in OCD patients yet, these results suggest that cortical measures could be helpful for better understanding the neural underpinnings of insight in OCD.

There are also potential links between subcortical volumes and insight. For example, verbal memory deficit was associated with poor insight in OCD (Kashyap et al., 2012; Kitis et al., 2007; Tumkaya et al., 2009), whose correlation may lie in regions beyond the cortex, especially in the hippocampus, which was generally regarded to be the core region for memory processing. Fan et al. (2017a) have found decreased spontaneous neuronal activity in the left hippocampus in OCD patients with poor insight when compared with healthy controls. Moreover, McFarland et al. (2013) detected a relationship between increased caudate, putamen and thalamus volumes, and impaired insight in first-episode affective and non-affective psychosis patients.

Thus, the present study was designed to clarify the neuroanatomical correlates of insight by comparing both the cortical architectures and subcortical volumes in OCD patients with good versus poor insight. As psychotropic medication and age are known to affect cortical structures in OCD (Boedhoe et al., 2017; Fouche et al., 2017), we conducted our analyses in a sample of drug-naïve adults. As a means of comparison, a healthy control group was also included. Although this study is primarily exploratory, we expected to replicate previous results of cortical abnormalities in multiple brain regions in both OCD groups, with structural changes in OCD patients with poor insight presenting in a more widespread and severe manner. We also expected to find a relationship between decreased cortical thickness in the frontal, parietal and temporal cortices, as well as decreased hippocampus volume, and poor insight in OCD.

2. Materials and methods

2.1. Participants

Forty-seven drug-naïve OCD patients were recruited from the psychology clinic at Second and Third Xiangya Hospital of Central South University. To be eligible, patients fulfilled criteria for OCD, as confirmed by the Structured Clinical Interview for DSM-IV-TR Axis I Disorders-Patient Edition (First et al., 2001), administered to each patient by two experienced clinical psychiatrists. Exclusion criteria included: (a) a diagnosis of other axis I and II psychiatric disorders; (b) a present or previous history of major medical or neurological problems; (c) a present or previous history of drug or alcohol abuse; (d) pregnancy; (e) unable to obtain MRI. Forty-two healthy people were recruited as healthy controls (HCs). Each of the HCs underwent the Structured Clinical Interview for DSM-IV-TR Axis I Disorders-Non-patient version (First et al., 2002) to exclude any current or past Axis I and II disorders. The other exclusion criteria were the same as in the patient group.

All subjects were right-handed as assessed by the Edinburgh Handedness Inventory (Oldfield, 1971) and have an education for more than 9 years. The study was approved by the Ethics Committee of the Second Xiangya Hospital of Central South University. Prior to participating in this study, all subjects signed a written informed consent after being told all procedures involved.

2.2. Clinical assessment

OC symptoms severity was assessed in patients using the Yale-Brown Obsessive-Compulsive Scale (Y-BOCS; Goodman et al., 1989). The Chinese version of the Brown Assessment of Beliefs Scale (BABS; Niu et al., 2016) was used to measure insight levels. It is a 7-item semi-structured scale that was developed to evaluate multidimensional construct of insight across a variety of psychiatric conditions. Each item is rated on a 5-point Likert scale, ranging from 0 (non-delusional or least pathological) to 4 (delusional or most pathological). Patients were divided into two groups based on total BABS score in conjunction with rating on Item #1 ‘conviction’: OCD patients with poor insight (OCD-PI; total BABS score ≥ 12 and Item #1 score ≥ 3) and patients with good insight (OCD-GI; total BABS score < 12 or Item #1 score < 3) (Fan et al., 2017a, 2017b). Consequently, of the 47 OCD patients enrolled, 21 were classified into the OCD-PI group and 26 into the OCD-GI group.

The Beck Depression Inventory (BDI; Beck et al., 1961) and the State-Trait Anxiety Inventory (STAI; Spielberger, 1983) were used to assess the severity of depressive and anxiety symptoms experienced by each participant, respectively, among all participants.

2.3. Image acquisition

MRI scans were collected on a Siemens Skyra 3-T magnetic resonance scanner with a 32-channel head coil at the Second Xiangya
Cortical surface reconstruction and volumetric segmentation were performed using the FreeSurfer software version 5.3.0 (http://surfer.nmr.mgh.harvard.edu/). The FreeSurfer automated processing pipeline was used, briefly involving motion correction, skull stripping, registration to FreeSurfer’s Talairach atlas, intensity normalization, subcortical segmentation and cortical parcellation based on the Desikan-Killiany Atlas (Desikan et al., 2006; Fischl and Dale, 2000; Fischl et al., 2004). Cortical surface reconstruction and volumetric segmentation were performed using the FreeSurfer software version 5.3.0 (http://surfer.nmr.mgh.harvard.edu/). The FreeSurfer automated processing pipeline was used, briefly involving motion correction, skull stripping, registration to FreeSurfer’s Talairach atlas, intensity normalization, subcortical segmentation and cortical parcellation based on the Desikan-Killiany Atlas (Desikan et al., 2006; Fischl and Dale, 2000; Fischl et al., 2004). Cortical thickness was defined as the distance between equivalent vertices of the grey/white boundary and pial surface, whereas surface area was calculated as the sum of the area of the vertices falling within a given region on the white matter surface. Volumes of seven subcortical regions encompassing accumbens, amygdala, caudate nucleus, hippocampus, pallidum, putamen and thalamus, and estimated intracranial volume (eICV) were obtained from the automated procedure for volumetric measures implemented in FreeSurfer (Fischl et al., 2002, 2004).

2.4. Image processing

OCD-GI, OCD with good insight; OCD-PI, OCD with poor insight; HCs, healthy controls; Y-BOCS, Yale-Brown Obsessive-Compulsive Scale; STAI-S, Spielberger State-Trait Anxiety Inventory-State Form; STAI-T, Spielberger State-Trait Anxiety Inventory-Trait Form; BDI, Beck Depression Inventory; BABS, Brown Assessment of Beliefs Scale.

F/t2χ2: variables of age, education, STAI-T, STAI-S, and BDI were tested by one-way ANOVAs (results were indicated by F); Categorical data such as gender was tested using chi-squared tests (results were indicated by χ2), and variables such as age of onset, duration, Y-BOCS and BABS were statistically tested by two-sample t-test (results were indicated by t); p, statistical significance, significant at p < 0.05.

Significant post hoc tests: STAI-S: OCD-GI > HCs, OCD-PI > HCs (p < 0.001); STAI-T: OCD-GI > HCs, OCD-PI > HCs (p < 0.001); BDI: OCD-GI > HCs, OCD-PI > HCs (p < 0.001).

Hospital of Central South University. For each subject, we obtained high resolution brain structural images by using a T1-weighted sagittal magnetization-prepared, rapid acquisition gradient-echo (MPRAGE) sequence. The sequence parameters were as follows: repetition time (TR) = 1900 ms, echo time (TE) = 2.01 ms, flip angle (FA) = 9°, field of view (FOV) = 256 mm, slice thickness = 1.0 mm, matrix = 256 × 256, voxel size = 1.0 × 1.0 × 1.0 mm³ and 176 slices covering the whole brain.

2.5. Statistical analysis

Statistical analyses were carried out using SPSS software version 18.0. Demographic and clinical variables were compared with χ2 test for categorical variables and independent-sample t-tests or one-way ANOVA for continuous variables. Pearson correlations were conducted to evaluate the relationships between insight levels and other clinical variables in OCD (e.g., Y-BOCS, BDI, and STAI scores, age of onset and illness duration).

Cortical measures were compared in a pairwise manner among groups by fitting a general linear model at each vertex with cortical thickness and cortical surface area respectively as dependent variables, group (OCD-PI, OCD-GI or HCs) as the independent variable, and age and gender as covariates. For the comparison between OCD-GI and OCD-PI, additional covariates including Y-BOCS and BDI scores, and age of onset were also controlled due to their potential effects on cortical measures (Boedhoe et al., 2017; Piras et al., 2015) and/or their relationship with the BABS score (see details in Results section). To this end, vertex-wise estimates of cortical surface architecture were mapped into FreeSurfer default template and smoothed using a Gaussian kernel with a full width at half maximum (FWHM) of 15 mm. All results were corrected for multiple comparisons using a cluster-wise Monte Carlo simulation implemented in FreeSurfer (Hagler et al., al. 2006; Hayasaka and Nichols 2003). The Monte Carlo simulations test the data against an empirical null distribution of maximal suprathreshold cluster size across 10,000 iterations, using a primary cluster-forming threshold of p < 0.05, yielding clusters fully corrected for multiple comparisons across the surfaces. The p-values of the resulting clusters of the original data expressed as cluster-wise probability (CWP) represent the probability of detecting a maximum cluster of that size or larger during the simulation step. Clusters with a CWP p < 0.05 were regarded as significant in this study. Only clusters that survived multiple corrections were presented in the Results.

For the subcortical analyses, we used ANCOVAs with each subcortical volume as the dependent variable, group (OCD-PI, OCD-GI or HCs) as the independent variable, and age, gender, BDI scores, and eICV as covariates. Significant results (p < 0.05) were corrected using Bonferroni-Holmes procedure.

To further examine core regions that could account for poor insight in OCD, mean values were extracted from those impaired regions where significant differences in cortical thickness and surface area were observed between OCD-PI and OCD-GI. Partial correlation analyses between mean cortical values and subcortical measures, and BABS scores were then examined in OCD-GI and OCD-PI group separately, with the Y-BOCS score controlled as a covariate. Multiple comparisons for these correlation analyses were corrected using Bonferroni-Holmes procedure.

3. Results

3.1. Demographic and clinical characteristics

Demographic and clinical characteristics of the participants are found in Table 1.

There were no significant group differences in terms of age, gender, and education levels. Subscales of STAI and BDI scores differed among the three groups, with both OCD-PI and OCD-GI groups showing higher scores than HCs on all three measures, and no difference observed between the patient groups. Furthermore, no significant difference was found between the OCD-GI and OCD-PI in terms of age of onset, illness duration, and Y-BOCS scores. Among all the clinical variables, the BABS...
3.2.1. Cortical thickness

3.2.2. Cortical surface area

3.3. Group comparisons in subcortical volumes

3.4. Correlations between cortical thickness and insight level

Table 2
Correlations between BABS scores and other clinical variables in OCD patients.

|          | Y-BOCS | STAI-S | STAI-T | BDI    | Age of onset | Illness duration |
|----------|--------|--------|--------|--------|--------------|------------------|
|          | \( r \) | \( p \) | \( r \) | \( p \) | \( r \) | \( p \) | \( r \) | \( p \) |
| BABS     | 0.413  | 0.004  | 0.069  | 0.649  | −0.020       | 0.893            | 0.095           | 0.531           | 0.038           | 0.802           | −0.048          | 0.751            |

Y-BOCS, Yale-Brown Obsessive-Compulsive Scale; STAI-S, Spielberger State-Trait Anxiety Inventory-State Form; STAI-T, Spielberger State-Trait Anxiety Inventory-Trait Form; BDI, Beck Depression Inventory; BABS, Brown Assessment of Beliefs Scale. \( r \), Pearson’s correlation coefficient; \( p \), statistical significance, significant at \( p < 0.05 \).

Fig. 1. (A) Brain regions exhibiting differences in cortical thickness between OCD-GI and HC. (B) Brain regions exhibiting differences in cortical thickness between OCD-PI and HC. (C) Brain regions exhibiting differences in cortical thickness between OCD-PI and OCD-GI. Results are shown at \( p < 0.05 \) corrected for multiple comparisons with Monte-Carlo simulation. Clusters color-coded in blue indicate significantly decreased cortical thickness in the OCD-PI group compared to either the OCD-GI or HCs or in the OCD-GI group compared to the HCs. Clusters are overlaid on average inflated images with sulci displayed as dark relative to gyri.

4. Discussion

To our knowledge, this is the first study to date having examined neurobiological basis underlying poor insight in OCD using an SBM approach, in which cortical thickness, cortical surface area as well as subcortical volumes were comprehensively compared among OCD patients with good and poor insight, and HCs. We observed a thinner cortex in OCD-PI in the left superior frontal gyrus and ACC as well as in the right lateral parietal region, when compared to both the OCD-GI group and HCs. Moreover, mean values of cortical thickness extracted from each cluster were further correlated with insight levels in the OCD-PI patients. By choosing a cohort of drug-naïve adult patients with OCD and controlling for illness severity, age of onset and depressive symptoms, we ruled out the effects of potential clinical confounders.

Consistent with our first hypothesis, we replicated previous findings.
by demonstrating decreased cortical thickness in the middle and inferior temporal gyri in both OCD-GI and OCD-PI (Fouche et al., 2017; Shin et al., 2007). Piras et al. (2015) also proposed a dorsolateral prefronto-striatal "executive'' circuit to explain the involvement of temporal region in the pathophysiology of OCD. Our results thus indicate that cortical thickness is a sensitive and solid enough index to understand the neurobiological mechanisms associated with structural abnormalities in OCD. On the other hand, OCD-PI and OCD-GI displayed some distinct cortical characteristics. For example, while OCD-PI showed cortical thinning in the temporal pole, the OCD-GI showed that in the superior temporal gyrus (STG). Even for the same structures, their specific locations were different. This finding could help to explain the inconsistency in previous results, i.e., while some studies showed cortical thinning in a certain area, such as STG, others not. Besides, we observed more widespread cortical abnormalities encompassing medial frontal, lateral parietal and occipital regions in OCD-PI, which support the notion that the specifier 'poor insight' helps to classify a subgroup of patients at the more severe end of OCD spectrum (Catapano et al., 2010).

The most outstanding finding of the current study is the identification of an association between cortical thinning in the left medial frontal cortex (including ACC) with poor insight. This decreased cortical thickness, which biologically represents the reduction of neurons within cortical columns, may be regarded as a reflection of insight-related neurodegenerative changes as having been suggested in a previous NSS study (Karadag et al., 2011). Since cortical thickness constitutes a reliable measure to identify brain-behavior relationships between regional structural changes and cognitive performance (Dickerson et al., 2008), and there is strong evidence of associations between reduced cortical thickness and impaired cognitive performance both in healthy population and disease (Hartberg et al., 2010; Tuladhar et al., 2015), we would also like to explain our result from a functional perspective, mainly on the basis of connections between the median walls, particularly the ACC and the medial prefrontal cortex (mPFC), with limbic structures in an effort to management and integrate sensory and emotional information (Ritkin et al., 2006; 2011). Some researchers have described appraisal functions of the mPFC and the ACC, such as representing the value of stimuli or actions (Kalisch et al., 2006; Ochsner and Gross, 2005; Rushworth et al., 2007). Decreased thickness of these areas may therefore be related to a deficit in the integration and evaluation of incoming signals from afferent limbic structures, which causes difficulty in tagging unusual mental events as pathological.

Besides, our results revealed that OCD-PI had decreased thickness in the right lateral parietal cortex, especially in the inferior parietal gyrus, compared with both OCD-PI and HCs. Moreover, in a poor insight cohort, there is a significant relationship between cortical thickness and insight level in this region. Our failure in detecting such a significant relationship in the OCD-GI group may be due to the relatively small sample size of the current study. Clearly, though, the parietal cortex played a role in determining insight in OCD patients. Interestingly, previous studies have linked lesions in the parietal lobe, especially in the right hemisphere, to anosognosia (i.e., unawareness of symptoms in neurological deficit, Bisiach et al., 1986; Ramachandran, 1995) and found reduced oligodendroglial density in the inferior parietal lobule (IPL) in schizophrenia patients with poor insight (Vostrikov et al., 2013). These similarities in insight-related structural abnormalities indicated that various psychiatric disorders and primary neurological disorders (associated with anosognosia) may share a similar basis of poor insight. This is also supported by evidence that frontal abnormalities have ascribed a key role in the pathogenesis of poor insight in psychosis (Flashment et al., 2001; Sapaea et al., 2007; Shad et al.,...
The involvement of the inferior parietal cortex may be explained by its role in self-other discrimination (Uddin et al., 2006). Specifically, a network compassing the mPFC, ACC and IPL, but also the PCC and insula has been proposed to be involved in the self-reflection (van der Meer et al., 2010), an ability to process information about the "self", and in patients, representing their capacity to self-reflect, to question unusual mental experiences and willingness to consider alternative explanations. Functional MRI studies have demonstrated an involvement of self-reflection networks underlying poor insight in schizophrenia (Curcić-Blake et al., 2015; van der Meer et al., 2012). Moreover, a specific metacognitive dimension of insight (i.e. cognitive insight) has been hypothesized (Orfei et al., 2013; Buchy et al., 2016) and in these studies, reduced volume or thickness in the prefrontal and parietal regions has been reported to correlate with lower self-reflectiveness. Hence, it is plausible that decreased thickness in the medial frontal and inferior parietal regions was involved in the pathophysiology of insight in OCD via their role in metacognition, especially in self-reflectiveness. This would be true since impaired metacognition and theory of mind (ToM) have been reported in OCD patients with poor insight (Önen et al., 2013; Tulaci et al., 2018). However, it still needs further functional MRI studies which include self-reflection tasks to confirm these associations and elucidate its neural mechanism.

Finally, we failed to find a significant relationship between subcortical volumes, for instance, the hippocampus volume, and insight, indicating that the effect of subcortical structures on insight may be minimal. This may be due to insight in OCD was related to higher-order cognitive processing such as executive function, verbal working memory and metacognition (Kashyap et al., 2012; Önen et al., 2013; Tumkaya et al., 2009), whose correlations are mainly lie in cortical structures. Another explanation is that the subcomponents of subcortical structures may have different roles. Further studies that assess the link between insight and subfields of these subcortical structures may draw out a different picture.

There are several limitations to be mentioned in the present study. First, it should be noted that the BABS employed in our study assessed the severity of a multidimensional construct of insight resulting in a total score. Such being this case, individuals who rate the same scores on this measure could exhibit very different insight profiles. Unfortunately, this problem has not been resolved, and the BABS is still one of the best and the most popular instruments available for assessing insight in OCD; second, this is a preliminary study restricted to structural neural correlates of insight with relatively small sample size. Future studies to replicate and extend these findings are needed.

5. Conclusions

Taken together, an association between thickness reduction in the left superior frontal gyrus, the left ACC and the right inferior parietal regions, and poor insight in OCD patients was observed. We thus propose that cortical thinning in the left medial frontal and right inferior parietal cortices may be neural underpinnings of poor insight in OCD. In spite of the aforementioned limitations, our structural data points to the need for future studies to investigate the structural, functional, and neurochemical correlates of insight deficits in OCD.

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CRediT authorship contribution statement

Wangting Liu: Formal analysis, Writing - original draft. Jun Gan: Data curation. Jie Fan: Data curation. Hong Zheng: Data curation. Sihui Li: Data curation. Raymond C.K. Chan: Supervision. Changlian Tan: Conceptualization. Xiongzhao Zhu: Conceptualization.

Declaration of Competing Interest

None.

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