Cortical thickness abnormalities in patients with first episode psychosis: a meta-analysis of psychoradiologic studies and replication in an independent sample

Keren Wen1,1, Youjin Zhao1,2,1, Qiyong Gong1,3,1, Qiyong Gong1,3,1, Ziyu Zhu1, Qian Li1, Nanfang Pan1, Shiqin Fu1, Joaquim Radua4,5,6, Eduard Vieta4,7, Poornima Kumar8,9, Graham J. Kemp10 and Bharat B. Biswal11,12

1Huaxi MR Research Center (HMRRC), Department of Radiology, West China Hospital of Sichuan University, Chengdu 610041, Sichuan, China
2Research Unit of Psychoradiology, Chinese Academy of Medical Sciences, Chengdu 610041, Sichuan, China
3Functional and Molecular Imaging Key Laboratory of Sichuan Province, Chengdu 610041, Sichuan, China
4Institut d’Investigacions Biomèdiques August Pi i Sunyer (IDIBAPS), Mental Health Research Networking Center (CIBERSAM), Barcelona 08036, Catalonia, Spain
5Centre for Psychiatric Research and Education, Department of Clinical Neuroscience, Karolinska Institutet, Solna 171-77, Stockholm, Sweden
6Department of Psychosis Studies, Institute of Psychiatry, Psychology and Neuroscience, King’s College London, London WC2R 2LS, UK
7Barcelona Bipolar Disorders and Depressive Unit, Hospital Clinic, Institute of Neurosciences, University of Barcelona, Barcelona 08036, Catalonia, Spain
8Center for Depression, Anxiety and Stress Research, McLean Hospital, Belmont 02478, MA, USA
9Department of Psychiatry, Harvard Medical School, Boston 02115, MA, USA
10Liverpool Magnetic Resonance Imaging Centre (LiMRIC) and Institute of Life Course and Medical Sciences, University of Liverpool, Liverpool L69 3GE, UK
11Department of Biomedical Engineering, New Jersey Institute of Technology, Newark 07102, NJ, USA
12The Clinical Hospital of Chengdu Brain Science Institute, MOE Key Laboratory for Neuroinformation, University of Electronic Science and Technology of China, Chengdu 610054, Sichuan, China

*Correspondence: Qiyong Gong, qiyonggong@hmrrc.org.cn
#Keren Wen and Youjin Zhao contributed to this work equally.
Abstract

Background: Abnormalities of cortical thickness (CTh) in patients with their first episode psychosis (FEP) have been frequently reported, but findings are inconsistent.

Objective: To define the most consistent CTh changes in patients with FEP by meta-analysis of published whole-brain studies.

Methods: The meta-analysis used seed-based d mapping (SDM) software to obtain the most prominent regional CTh changes in FEP, and meta-regression analyses to explore the effects of demographics and clinical characteristics. The meta-analysis results were verified in an independent sample of 142 FEP patients and 142 age- and sex-matched healthy controls (HCs), using both a vertex-wise and a region of interest analysis, with multiple comparisons correction.

Results: The meta-analysis identified lower CTh in the right middle temporal cortex (MTC) extending to superior temporal cortex (STC), insula, and anterior cingulate cortex (ACC) in FEP compared with HCs. No significant correlations were identified between CTh alterations and demographic or clinical variables. These results were replicated in the independent dataset analysis.

Conclusion: This study identifies a robust pattern of cortical abnormalities in FEP and extends understanding of gray matter abnormalities and pathological mechanisms in FEP.

Key words: psychoradiology; first episode psychosis; early psychosis; cortical thickness; meta-analysis; seed-based d mapping

Introduction

First episode psychosis (FEP) refers to the initial stage of psychosis (Breitborde et al., 2009), usually characterized by distortions in cognition, behavior, and physical function (Chudleigh et al., 2011; Grant et al., 2001). FEP is common in young adults (Reed, 2008) and has high rates of disability and mortality (Simon et al., 2018). With the development of psychoradiology (Gong, 2020; Lui et al., 2016), an abundance of structural magnetic resonance imaging (MRI) studies have provided key information for neurostructural changes in FEP (Li et al., 2021). Applying surface-based morphometric methods allows quantification of cortical thickness (CTh) (Fischl and Dale, 2000). CTh reflects the density, arrangement, and size of neurons, neuroglia, and nerve fibers (Narr et al., 2005), and is only minimally affected by partial volume effects (Winkler et al., 2010). CTh abnormalities reflect regional disease-specific effects: cortical thinning can follow a loss of dendrites and dendritic spines or alterations during myelination, while neuroinflammation or other factors can increase CTh (Hutton et al., 2008; Narr et al., 2005; Sowell et al., 2004; Winkler et al., 2010).

Recent studies in FEP have reported extensive CTh alterations in salience processing, cognitive, and emotional areas involving the default mode network (DMN) and the salience network (SN). However, reported results are inconsistent: compared with healthy controls (HCs), lower CTh has been reported in the superior (Ansell et al., 2015; Haring et al., 2016; Plitman et al., 2016), middle (Ansell et al., 2015; Buchy et al., 2017; Plitman et al., 2016), and inferior (Plitman et al., 2016; Qiu et al., 2013) frontal gyri, superior (Plitman et al., 2016; Schultz et al., 2010) and inferior parietal (Ansell et al., 2015; Bodnar et al., 2014) gyri; superior (Scanlon et al., 2014; Schultz et al., 2010; Song et al., 2015) and middle temporal gyri (Kim et al., 2012; Qiu et al., 2013; Scanlon et al., 2014); insula (Song et al., 2015); and cingulate gyri (Bodnar et al., 2014; Schultz et al., 2010). Higher CTh has been reported in the middle temporal gyrus (Qiu et al., 2013), temporal pole (Haring et al., 2016; Xia et al., 2015), precentral (Ansell et al., 2015; Haring et al., 2016), and postcentral gyri (Ansell et al., 2015), and bilateral superior, middle, and inferior occipital gyri (Ansell et al., 2015; Dukart et al., 2017). Some studies report no significant differences of CTh (Lesh et al., 2015; Lin et al., 2019; Reniers et al., 2015). Causes of these inconsistencies may include differences in sample characteristics (e.g. patient demographics, symptom severity, illness duration, medication status, and sample size), MRI data acquisition methods, and processing protocols. A meta-analysis is urgently needed to resolve these apparent contradictions. For brain imaging studies, coordinate-based meta-analysis can synthesize results from different studies in an unbiased way, yielding a robust picture of cortical alterations in disease (Quah and Cockerham, 2017).

Therefore, we aimed to conduct a meta-analysis of whole brain CTh studies in patients with FEP by using seed-based d mapping (SDM) software. We used a mask specifically designed for whole-brain CTh, which minimizes the effect of subcortical gray matter (Li et al., 2020). We performed meta-regression analyses to explore the effects of demographics and clinical characteristics on CTh. We also investigated whether the CTh findings in our meta-analysis could be replicated in an independent FEP sample: we used both a whole-brain vertex-wise analysis, and also (to directly compare cerebral areas identified in the meta-analysis) a region of interest (ROI)-based analysis. As previous studies report widespread disruptions of CTh in the DMN and SN, we hypothesized that, compared with HCs, FEP patients would show CTh...
alterations in DMN and SN involving core regions for cognitive and emotional function, possibly associated with demographics and clinical characteristics.

Materials and Methods
Meta-analysis of cortical thickness studies

Study selection
A comprehensive strategy was executed in September 2020 to search for pertinent literature in Web of Science, PubMed, Embase, and Science Direct databases. The search terms included: (“first-episode psychosis” OR “first-episode schizophrenia” OR “first-episode bipolar disorder” OR “first-episode depression” OR “early psychosis” OR “early schizophrenia” OR “early bipolar disorder” OR “early depression” OR “drug-naïve psychosis” OR “drug-naïve schizophrenia” OR “drug-naïve bipolar disorder” OR “drug-naïve depression”) AND (“cortical thickness”). In addition, we manually checked the references of these studies to identify further studies for inclusion. Studies meeting the following criteria were included: (i) studies published in English in a peer-reviewed journal; (ii) studies that recruited individuals who met the diagnostic criteria of FEP and healthy controls based on the cross-diagnostic approach (Fusar-Poli and Meyer-Lindenberg, 2013a; Fusar-Poli and Meyer-Lindenberg, 2013b), where FEP was defined as several mental illnesses characterized by psychotic symptoms including schizophrenia spectrum psychoses (schizophrenia, schizoaffective, schizophreniform) and affective psychoses (bipolar psychosis and psychotic depression); (iii) studies that used whole-brain CTh analysis methods; and (iv) studies that documented Montreal Neurological Institute (MNI) or Talairach (TAL) coordinates of peak CTh alterations, or reported no significant findings. Reviews and theoretical papers were excluded. For studies using overlapping samples, we only included the study with largest samples. For longitudinal studies, we only included baseline data to avoid bias toward the effect of interventions or illness progression. For studies containing multiple independent patient samples, results were considered as separate datasets. Corresponding authors were contacted if the peak coordinates of effects were not available in whole-brain CTh studies. Two authors independently conducted the literature search, and any inconsistencies were discussed to reach an agreement. The flow of literature selection is summarized in Supplementary Fig. S1.

Quality assessment and data recording
Primary studies included in meta-analysis were assessed using a modified version of Newcastle-Ottawa Scale (NOS) that scores potential risk of bias for case-control studies in terms of six aspects of study quality (Keramatian et al., 2021; Wells et al., 2000). The details are given in Supplementary Table S1. The recorded information for each included dataset consisted of clinical characteristics (sample numbers, gender, mean age of participants, mean age at onset, mean illness duration, mean duration of untreated psychosis (DUP), mean symptom severity, medication status, diagnosis), imaging characteristics (scanner manufacturer and model, field strength, sequence name, spatial resolution, normalization template, repetition time/echo time, data processing software, analytic model, method to correct whole-brain results for multiple comparisons, and statistical threshold of the main findings), and the main CTh alterations. Peak coordinates and corresponding t, P, or z values were also recorded for SDM calculations. If only P or z values were available, these were converted into t values (O’Neill et al., 2019).

Protocol for meta-analysis
First, we performed a pooled meta-analysis with all included studies. Then subgroup meta-analyses of studies using 3.0 T and 1.5 T MRI scanners were performed to investigate the possible effects of field strength. SDM (www.sdmproject.com) software (Kimmel et al., 2016; Zhang et al., 2016) was used to execute the meta-analysis. The details have been presented in detail elsewhere (Radua et al., 2011; Radua and Mataix-Cols, 2009; Radua et al., 2014b), and we give only a brief summary here. First, reported peak coordinates and t values of significant differences between FEP patients and HCs were used to recreate an effect-size signed map for each study. Importantly, applying the same threshold throughout the whole brain in each included study avoided biases toward brain regions with liberal thresholds (Radua et al., 2011; Radua et al., 2014b). Second, we chose a specially constructed mask (the details of which can be found in (Li et al., 2020)) to restrict the analysis to the cortex. Third, the mean map was derived using a traditional random-effects meta-analytic methods, with both negative and positive changes presented in the same map (Radua and Mataix-Cols, 2009). We used SDM’s default threshold for analyses (voxel P < 0.005, peak height z > 1, cluster extent = 10 voxels).

Jackknife, heterogeneity and publication bias analysis
Replicability was assessed by jackknife sensitivity analysis. The main statistical analysis was repeated N times (N = number of datasets in the meta-analysis) by discarding one dataset at a time to determine whether the results remained significant. We estimated the statistical (between-studies) heterogeneity of individual clusters using q statistics ($\chi^2$ distribution converted to z values) and tested for heterogeneity of findings with a permutation approach. The possibility of publication bias for regions showing altered CTh was examined using Egger tests.

Meta-regression analysis
Meta-regression analyses were conducted to examine the effects of age, illness duration (mean values), gender
(percentage of females), and medication status (percentage of medicated participants) on CTh. Meta-regression analysis of age at onset and symptom severity was not possible, being reported in fewer than nine studies (Radua and Mataix-Cols, 2009). In accordance with our previous meta-analysis, the probability threshold was reduced to 0.0005 to minimize detection of spurious relationships (Radua et al., 2011; Radua and Mataix-Cols, 2009), we required abnormalities to be detected both in the slope and in one of the extremes of the regressor, any regions not detected in the main analyses were discarded, and regression plots were visually inspected to discard findings driven by too few studies (Radua and Mataix-Cols, 2009).

Validation of meta-analysis results in an independent sample of patients and controls

To verify the results from the meta-analysis, a study was conducted using both whole-brain vertex-wise and ROI-based analyses to compare CTh between an independent group of FEP patients and controls.

Participants and MRI data acquisition

We recruited 142 patients with FEP and 142 age- and sex-matched HCs, and acquired high-resolution T1-weighted images. The severity of clinical symptoms and psychosocial functioning were evaluated using the Positive and Negative Syndrome Scale (PANSS) (Kay et al., 1987) and the Global Assessment Function (GAF) scale (Morosini et al., 2000). The inclusion and exclusion criteria and MRI acquisition parameters are given in the Supplementary Methods. This study was approved by the ethics committee of Sichuan University, China. Written informed consent was obtained from each participant.

Image processing

The processing of structural images was performed in FreeSurfer software package 6.0.0 (Fischl, 2012) (http://surfer.nmr.mgh.harvard.edu) using the following steps: (i) head motion correction, (ii) skull stripping, (iii) transformation to the Talairach space, (iv) segmentation of gray/white matter, (v) surface inflation and registration to a spherical atlas, (vi) CTh calculation as the shortest distance between the gray–white interface and the pial interface (Poppa and Bechara, 2018), and (vii) surface-map smoothing using a Gaussian kernel with a full-width at half-maximum of 20 mm.

Statistical analysis

As a preliminary analysis, Kolmogorov–Smirnov (KS) tests, Shapiro–Wilk (SW) tests, and visual assessment of the histogram were comprehensively used to analyze the normality of clinical variables (Ghasemi and Zahediasl, 2012). For normally distributed continuous variables, two-sample t-tests were used. For nonnormally distributed continuous variables, Mann–Whitney tests were used. For gender distribution, the chi-square test was used. All tests were two-tailed, and statistical significance was considered at P < 0.05.

In the independent sample, whole-brain vertex-wise analysis assessed the group differences of CTh using a general linear model (GLM) based on QDEC (query, design, estimate, contrast) in FreeSurfer. Monte Carlo simulation controlled for multiple comparisons (cluster-wise corrected P < 0.05). Correlations between CTh and age, GAF, and PANSS scores were separately examined for each vertex in the FEP group. Statistical significance of correlations was set at P < 0.05 after correction for multiple comparisons using the criteria of false discovery rate (FDR).

Results

The characteristics of the included studies

The meta-analysis incorporated 10 studies, with a total sample of 624 patients with FEP (213 females (34%), mean age 24.2 years), and 505 healthy controls (193 females (38%), mean age 24.5 years). Three of the 10 studies recruited two subgroups depending on the medication type (Ansell et al., 2015), medication status (Lesh et al., 2015) and history of cannabis use (Rais et al., 2010): Ansell et al. separated 52 FEP patients into two medication-type datasets: first-generation antipsychotics (FGA) and second-generation antipsychotics (SGA) included here as two datasets (Ansell et al., 2015). Lesh et al. studied two subgroups, unmedicated and medicated. We included only the former, as peak coordinates of CTh alterations were not available for the latter (Lesh et al., 2015). Rais et al. included two subgroups, cannabis-using and non-using (Rais et al., 2010); to reduce confounding factors we only included the latter. Thus, there were 11 datasets from 10 studies in the meta-analysis, and no significant differences were found in mean age (t = -0.298, P = 0.783) and gender (χ² = 1.745, P = 0.187) between FEP patients and HCs. The clinical characteristics of included datasets are listed in Table 1, and the imaging characteristics and
Table 1: The clinical characteristics of the 11 datasets included.

| Paper                              | Numbers (female) | Age (y) | Age at onset (y) | Illness duration | Symptom severity               | Medication status | Diagnosis: numbers | Main findings                                                                 |
|------------------------------------|------------------|---------|------------------|------------------|-------------------------------|-------------------|-------------------|-----------------------------------------------------------------------------|
| Ansell et al., 2015                |                  |         |                  |                  |                               |                   |                   | cortical thinning: right rostral, middle frontal and superior frontal gyr   |
|                                    | 25 (8)           | 28 (11) | 21.9             | 21.1             | NA                            | 42 d              | SCZ: 15; depression disorders and BD: 10                                  |
|                                    |                  |         |                  |                  | PANSS: 22.4 (pos), 22.0 (neg), |                   | 25/161/39          | cortical thinning: right inferior parietal, superior frontal, and fusiform  |
|                                    |                  |         |                  |                  | 41.6 (gen)                     |                   |                  | gyri; cortical thickening: left precentral and right postcentral gyri       |
|                                    |                  |         |                  |                  |                               |                   |                  | No significant CTh alteration by whole-brain vertex-wise method             |
| Ansell et al., 2015                | 27 (9)           | 28 (11) | 21.9             | 21.1             | NA                            | 93 d              | SCZ: 12; depression disorders and BD: 15                                  |
|                                    |                  |         |                  |                  | PANSS: 22.4 (pos), 21.9 (neg), |                   | 27/252/165          | cortical thickening: left superior, middle and inferior occipital gyri,    |
|                                    |                  |         |                  |                  | 44.3 (gen)                     |                   |                  | fusiform gyr, occipital pole, fusiform and lingual gyrus, calcarene cortex |
| Buchy et al., 2018                 | 130 (37)         | 52 (15) | 24.1             | 24.3             | NA                            | 5.8 y             | 130/793/NA          | cortical thickening: left superior, middle and inferior occipital gyri,    |
|                                    |                  |         |                  |                  | SAPS: 4.0 (total pos), 1.5 (delusions); SANS: 8.8; Self-reflectiveness: 12.4; Self-certainty: 7.9; CDSS: 2.3 |                   |                  | fusiform gyr, occipital pole, fusiform and lingual gyrus, calcarene cortex |
|                                    |                  |         |                  |                  |                               |                   |                  | No significant CTh alteration by whole-brain vertex-wise method             |
| Dukart et al., 2017                | 59 (17)          | 26 (14) | 26.4             | 27.7             | NA                            | NA                | 28/216/NA          | cortical thickening: left superior, middle and inferior occipital gyri,    |
|                                    |                  |         |                  |                  | SANS: 18.0; BPRS: 49.7         |                   |                  | fusiform gyr, occipital pole, fusiform and lingual gyrus, calcarene cortex |
|                                    |                  |         |                  |                  |                               |                   |                  | No significant CTh alteration by whole-brain vertex-wise method             |
| Gutierrez et al., 2010             | 37 (12)          | 38 (16) | 26.8             | 25.0             | NA                            | 10.4 m            | 37/NA/80           | cortical thickening: left superior, middle and inferior occipital gyri,    |
|                                    |                  |         |                  |                  | WMS: 5.5; Planning: 7.2; WMM: 33.9; RAVLT: 38 |                   |                  | fusiform gyr, occipital pole, cerebellum exterior, right superior, middle and inferior occipital gyri, fusiform and lingual gyrus, calcarene cortex |
|                                    |                  |         |                  |                  |                               |                   |                  | SCZ: 34; BD: 1; depressed subtypes: 2                                     |
|                                    |                  |         |                  |                  |                               |                   |                  | No significant CTh alteration by whole-brain vertex-wise method             |
| Paper                        | Numbers (female) | Age (y) | Age at onset (y) | Illness duration | Symptom severity | Medication status | Diagnosis: numbers | Main findings                                      |
|------------------------------|------------------|---------|------------------|------------------|------------------|------------------|-------------------|----------------------------------------------------|
| Haukvik et al., 2016         | 79 (27)          | 82 (28) | 27.6             | 29.3             | 23.8             | 123 w            | NA                | SCZ: 37; SCZ-F: 2; SAD: 5; BD: 18; psychotic depression: 5; paranoid psychosis: 2; psychosis NOS: 10 |
| Lesh et al., 2015 b          | 22 (3)           | 37 (10) | 20.2             | 19.7             | NA               | 210 d            | 0/0/0             | SCZ/SAD/SCZ-F: 22 No significant CTh alteration in unmedicated group compared with HC |
| Lin, et al., 2019            | 145 (76)         | 147 (76) | 24.5             | 25.9             | 23.6             | 11 m             | 0/0/0             | SCZ/SCZ-F: 146 No significant CTh alteration by whole-brain vertex-wise method |
| Rais et al., 2010 c          | 32 (6)           | 31 (6)  | 23.3             | 24.7             | 22.5             | 351 d            | 18/NA/77          | SCZ: 32 No significant CTh alteration in non-cannabis-using group compared with HC |
| Reniers et al., 2015         | 22 (4)           | 22 (4)  | 20.6             | 22.5             | NA               | 3 w              | 14/200/5          | SCZ: 1; SAD: 3; SCZ-F: 9; SIFD: 2; psychosis NOS: 2; DD: 1; MDD: 2; bipolar unspecified: 1; BD: 1 No significant CTh alteration by whole-brain vertex-wise method |
Table 1: Continued

| Paper | Numbers (female) | Age (y) | Illness duration (y) | Symptom severity | Diagnosis: numbers | Main findings | Medication status d |
|-------|-----------------|---------|----------------------|-----------------|-------------------|--------------|-------------------|
| Scalon et al., 2014 | 46 (14) | 28.6 | 14 m | PANSS: 17 (pos), 15 (neg), 33 (gen); GAF: 51 | SCZ: 15; SAD: 4; DD: 3; mania: 9; psychotic: 6; psychosisNOS: 4 | Cortical thinning: right superior temporal gyrus and sulcus, extending into middle temporal gyrus | 3P2C | 22/18 |

Assessment of risk of bias

The potential risk of bias of the studies is described in Supplementary Figs S2 and S3. Four (40%) of the 10 studies included in the meta-analysis received a maximum score of 6 on the modified version of the NOS, five studies (50%) received a score of 5, and one study (10%) received a score of 4. All 10 studies are relatively low risk or with some concerns about bias, mostly because of no description about the selection of controls.

Results of the pooled meta-analysis

Compared with HCs, FEP patients showed lower CTh in the right middle temporal cortex (MTC) extending to superior temporal cortex (STC), insula, and anterior cingulate cortex (ACC) (Table 2 and Fig. 1).

Jackknife, heterogeneity, and publication bias

In whole-brain jackknife sensitivity analysis, lower CTh in the right MTC, insula, and ACC was highly replicable, being statistically significant in all but one combination (Supplementary Table S3). No statistically significant heterogeneity was detected. No statistically significant publication bias (Supplementary Fig. S4) was revealed in the right MTC ($P = 0.662$), insula ($P = 0.188$), and ACC ($P = 0.243$).

Subgroup meta-analysis

For the eight datasets that used a 1.5 T scanner, FEP patients showed lower CTh in the right MTC extending to STC, insula, ACC, and inferior frontal cortex, and higher CTh in the left precentral cortex than HCs (Supplementary Table S4). For the three datasets that used a 3.0 T scanner, there were no significant differences between the two groups.

Meta-regression analysis

Meta-regression analysis revealed no statistically significant correlation between CTh and age, gender, illness duration, or medication status (Supplementary Table S5).

Validation of meta-analysis results in an independent sample

The studied FEP patients and HCs did not significantly differ in age or gender. Table 3 summarizes their clinical and demographic characteristics. In the whole-brain analysis, FEP patients compared to HC showed significantly lower CTh in the bilateral STC (extending to MTC, temporal pole, insula, and fusiform) and superior frontal main CTh alterations are listed in Supplementary Table S2.
Table 2: Three regions with decreased cortical thickness in patients with FEP compared to healthy controls.

| Brain regions                          | MNI coordinates | Effect size | Variance |
|----------------------------------------|-----------------|-------------|----------|
| Right middle temporal cortex           | 50, -20, -12    | −1.023      | 0.00136  |
| Right superior temporal cortex         | 40, 2, 14       | −1.136      | 0.00051  |
| Right insular cortex                   | 40, -20, -12    | −1.011      | 0.00175  |
| Right anterior cingulate cortex        | 14, 32, 28      | −0.101      | 0.010    |

Cluster breakdown (voxels, n)

- Right middle temporal cortex, BA 20, 21, 22, 48 (135)
- Right superior temporal cortex, BA 21, 22, 23, 48 (129)
- Right insula cortex, BA 48 (96)
- Right Rolandic operculum, BA 44 (65)
- Right anterior cingulate/pars cingulate cortex, BA 32 (53)

Figure 1: The areas and corresponding Montreal Neurological Institute (MNI) coordinates of decreased (blue) cortical thickness in participants with FEP compared with healthy controls in the pooled meta-analysis.

cortex (extending to anterior and posterior cingulate cortex), and higher CTh in the bilateral lingual, cuneus gyri, and right prefrontal cortex (Table 4 and Fig. 2). In the ROI analysis, FEP patients showed significantly lower CTh in the MTC, insula, and ACC (Supplementary Table S6). There was no significant association between CTh and clinical characteristics in either whole-brain or ROI analyses (Supplementary Table S7).

Discussion

We used a coordinate based meta-analysis approach to define the most robust cortical thickness alterations in patients with FEP, and to assess the effects of clinical and demographic factors. We found three regions within the default mode network and salience network that showed lower CTh (i.e. cortical thinning): the right middle temporal cortex extending to superior temporal cortex, insula, and anterior cingulate cortex. Previous meta-analyses have reported lower gray matter volume in the STC, insula, and ACC in FEP, which may be part of its neural substrate (Fusar-Poli et al., 2014; Radua et al., 2012a; Shah et al., 2017). There was no significant effect of age, gender, illness duration, or medication status on CTh in any of the three identified clusters.

Replication in an independent sample is a particular strength of this study. Consistent with the meta-analysis, the whole-brain analysis showed qualitatively similar abnormalities of clusters located in the MTC, STC, insula, and ACC. The ROI analysis also supported these
Table 3: Characteristics of the independent sample of patients with FEP and healthy controls.

| Characteristic          | FEP (N = 142) | HCs (N = 142) | P     |
|-------------------------|---------------|---------------|-------|
| Age (years)             | 25.2          | 26.4          | 0.077 |
| Gender (male/female)    | 61/81         | 61/81         | 0.001 |
| GAF                     | 29.9          | 10.6          |       |
| PANSS Total             | 89.7          | -             |       |
| Positive                | 24.9          | 6.4           |       |
| Negative                | 19.1          | 8.3           |       |
| General psychopathology | 45.7          | 9.7           |       |

P by Mann–Whitney test
P by two-tailed Pearson chi-square test. Abbreviations: GAF, Global Assessment of Functioning Scale; N, number of participants; PANSS, Positive and Negative Syndrome Scale; SD, standard deviation.

results. Interestingly, we found no significant associations between clinical symptom severity and CTh, suggesting that cortical thinning may be a trait marker of predisposition to FEP rather than a manifestation of the disease (Brent et al., 2013). Participants with psychotic experiences or at genetic/clinical high risk for developing psychiatric disorders show a decline in gray matter in the temporal, frontal, and cingulate cortices, which persisted in those who transformed to psychosis (Merritt et al., 2021). Consistent with this, cortical thinning in the STC, ACC, and insula has been found in both psychotic patients and their healthy first-degree relatives (Oertel-Knöchel et al., 2013), suggesting that these CTh reductions might represent an endophenotype of psychosis.

The most prominent cortical thinning in FEP was in the temporal cortex, especially STC and MTC. This region is located in the lateral portion of temporal cortex, and is a component of the DMN (Mulders et al., 2015). The DMN is crucial in thought processes, autobiographical memory, and conceiving others’ perspectives (Buckner et al., 2008), and dysfunction of the DMN, particularly the STC/MTC, is related to core symptoms of psychosis (Mallikarjun et al., 2018; O’Neill et al., 2019). The STC is thought to contain the auditory association cortical areas (Sun et al., 2009) and is a potential candidate for the neural basis of auditory hallucinations in psychosis (Allen et al., 2008). The STC also plays an important role in cognitive impairment and disorganized behavior in psychosis, which is related to loss of CTh (Kim et al., 2012; Walton et al., 2017). The MTC, particularly the middle part delineated in our study, plays an important role in semantic memory (Chang et al., 2011; Squire, 1992; Xu et al., 2015). Semantic memory dysfunction manifests as loss of general knowledge and information (Hui et al., 2012; Tan et al., 2020) increasingly considered important in psychosis because semantic memory deficits are related to the severity of core symptoms, including delusions (Rossell et al., 1999) and disorganized thought (Tan and Rossell, 2014).

We also found lower CTh in the insula and ACC, which are important hubs of the SN (Mulders et al., 2015). A review has identified the cardinal role of the SN in early-stage psychosis (Palaniyappan and Liddle, 2012b). The SN is essential in detection and integration of emotional and sensory stimuli (Downar et al., 2000). Misattribution of salience to external and internal stimuli has been proposed as a core feature of psychosis, underlining psychotic symptoms such as delusions and hallucinations (Palaniyappan and Liddle, 2012b). The insular cortex has extensive interconnectivity with many cortical areas and the limbic system (Jang et al., 2006). Insula dysfunction may lead to abnormalities in processes related to salience processing (Menon and Uddin, 2010), emotional appraisal, and social cognition (Eckert et al., 2009) that are characteristic of psychosis (Wylie and Tregellas, 2010). ROI analyses support insular cortical thinning in FEP (Roiz-Santianez et al., 2010; Wannan et al., 2019). Possible mechanisms include anomalies in cortical maturational processes that lead to loss of CTh (Kim et al., 2012; Todtenkopf et al., 2005). Dysfunction of ACC is well characterized for negative symptoms in psychosis, and ACC thinning might be involved in the increasing social withdrawal, which is a characteristic of the psychosis prodrome (Bersani et al., 2014). In support of this, FEP patients with persistent negative symptoms (PNS) showed a thinner cortex in the right ACC compared with HCs and FEP patients without PNS (Bohdan et al., 2014).

In subgroup analysis of studies done on 1.5 T scanners, results remained largely consistent with the pooled meta-analysis, with one additional significantly lower-CTh cluster in the right inferior frontal cortex and one higher-CTh cluster in the left precentral cortex. Since 8 out of 11 of the included datasets were at 1.5 T, this concordance is reasonable. The lack of similar results in the 3 T subgroup may due to the limited number of datasets (3/11 of the included datasets). The results of subgroup analyses should be regarded with caution: the minimum number of studies recommended...
Table 4: Significant changes in cortical thickness in patients with FEP compared to healthy controls in the independent sample.

| Brain region                  | Talairach coordinates (peak) | x     | y     | z       | P      | Size (mm²) | Cluster breakdown                                      |
|-------------------------------|------------------------------|-------|-------|---------|--------|------------|-------------------------------------------------------|
| **Individuals with FEP < healthy controls** |                              |       |       |         |        |            |                                                       |
| Right superior temporal cortex |                             | 40.1  | 1.9   | 20.9    | 0.0001 | 10 842     | Superior temporal cortex                               |
|                               |                              |       |       |         |        |            | Middle temporal cortex                                 |
|                               |                              |       |       |         |        |            | Insula                                                |
|                               |                              |       |       |         |        |            | Inferior temporal cortex                               |
|                               |                              |       |       |         |        |            | Fusiform                                              |
|                               |                              |       |       |         |        |            |                                                       |
| Right superior frontal cortex  |                             | 11.4  | 6.1   | 41.8    | 0.0001 | 4823       | Superior frontal cortex                                |
|                               |                              |       |       |         |        |            | Dorsal anterior cingulate cortex                       |
|                               |                              |       |       |         |        |            | Posterior cingulate cortex                             |
|                               |                              |       |       |         |        |            |                                                       |
| Left superior temporal cortex  |                             | −17.7 | 31.8  | −59.3   | 0.0001 | 15 417     | Superior temporal cortex                               |
|                               |                              |       |       |         |        |            | Middle temporal cortex                                 |
|                               |                              |       |       |         |        |            | Insula                                                |
|                               |                              |       |       |         |        |            | Inferior temporal cortex                               |
|                               |                              |       |       |         |        |            | Fusiform                                              |
|                               |                              |       |       |         |        |            |                                                       |
| Left superior frontal cortex   |                             | −10.6 | 0.7   | 41.8    | 0.0001 | 8992       | Superior frontal cortex                                |
|                               |                              |       |       |         |        |            | Dorsal anterior cingulate cortex                       |
|                               |                              |       |       |         |        |            | Posterior cingulate cortex                             |
|                               |                              |       |       |         |        |            | Precentral cortex                                      |
| **Individuals with FEP > healthy controls** |                           |       |       |         |        |            |                                                       |
| Right medial occipital cortex  |                             | 10.2  | −69.6 | 17.1    | 0.0001 | 5767       | Lingual cortex                                         |
|                               |                              |       |       |         |        |            | Cuneus cortex                                          |
| Right prefrontal cortex        |                             | 9.4   | 52.5  | −5.9    | 0.0001 | 5722       | Prefrontal cortex                                      |
| Left medial occipital cortex   |                             | −6.9  | −80.6 | 5.2     | 0.0001 | 4704       | Lingual gyrus                                          |
|                               |                              |       |       |         |        |            | Cuneus gyrus                                           |
Cortical thickness abnormalities in FEP

Figure 2: The areas showing lower (blue) and higher (orange) cortical thickness in participants with FEP compared with healthy control participants in the independent sample analysis.

for subgroup analyses in SDM software is 10 (Radua et al., 2014a). Future studies with larger samples and consistent field strength are needed to confirm this finding.

We further propose that the aforementioned brain regions do not affect the disease individually, but as an integration. For example, patients with FEP showed decreased integrity in the white matter fibers (such as cingulum) connecting the cingulate cortex and temporal areas (Sun et al., 2015), which may be a cause or a consequence of abnormalities in the gray matter (Konrad and Winterr, 2008). In terms of brain function, a recent meta-analysis comparing FEP with HCs reports hypo-connectivity between SN (insula and ACC) and DMN (particularly MTC), associated with perception anomalies (O’Neill et al., 2019). In addition, a multimodal meta-analysis has identified coupling changes between cortical structure (lower gray matter volume) and function (hyper-activation or hypo-activation) in these regions, and suggested a causal link: hyper-activation may cause gray matter loss by “exhaustion,” while hypo-activation may cause the same as a compensation to avoid neuronal underemployment (Radua et al., 2012a). Our meta-analysis adds information on cortical thinning.

Interestingly, cortical thinning in our meta-analysis only presented on the right hemisphere, which is consistent with the theory of psychosis as a disturbance of lateralization (Crow et al., 2013). In fact, asymmetry is ubiquitous in the human brain (de Kovel et al., 2019). An older study suggested that auditory hallucinations, the core feature of psychosis, arise from the right hemisphere, and perhaps because the lack of characteristic of being self-generated (Nasrallah, 1985). The right hemisphere is important in mediating higher language functions, of which the deficits displayed by psychosis patients may make a significant contribution to their social interaction deficits (Mitchell and Crow, 2005). Other psychotic symptoms of FEP, such as lack of insight and suicidal behavior, were linked with unilateral gray matter reduction in the right hemisphere (Canal-Rivero et al., 2020; Tordesillas-Gutierrez et al., 2018). Moreover, factors such as handedness, sex, and disease processes associated with psychosis have been suggested to modulate the structural lateralization of the cerebral hemispheres (Hamilton et al., 2007). We deduced that the right hemisphere lateralization of cortical thickness thinning was a combined effect of multiple causes. Future studies that will clarify the specific contribution of the right hemisphere to psychopathological mechanisms are expected.

The present study, together with previous findings of structural and functional abnormalities in insula, ACC, and MTC in FEP (Cui et al., 2018; Palaniyappan and Liddle, 2012a; Radua et al., 2012a), strongly implicates the SN and DMN in the pathogenesis of FEP.

Limitations

Our study has some limitations. A general limitation of coordinate-based meta-analysis is that using published coordinates instead of t-statistic brain maps limits accuracy (Radua et al., 2012b). Second, given the mixed clinical and imaging characteristics of our samples and the lack of some key information (e.g. diagnosis type, medication type and dosage, spatial resolution), we were unable to make much use of subgroup or regression analyses. Third, our study was restricted to cortical areas, and subcortical regions such as hippocampus, amygdala, and thalamus remain to be explored. Fourth, group-level inferences employing traditional mass-univariate neuroimaging approaches in our study restricted the information to make diagnostic decisions about FEP patients (Vieira et al., 2020).
Conclusion

In summary, in a whole-brain meta-analysis in FEP, we found prominent cortical thinning involving the DMN (i.e. MTC and STC) and SN (i.e. insula and ACC), which was replicated in an independent sample of patients and control participants. We suggest that these reflect the neuropathologic basis of deficits of salience processing, and cognitive and emotional integration, which are important in FEP. Future studies with medication-naïve FEP participants will help verify our conclusions. Longitudinal studies will provide crucial information in understanding the clinical progress of CTh alterations from high psychosis risk to chronic status.

Supplementary Data

Supplementary data are available at Psychoradiology online.

Author contributions

Qiyong Gong conceptualized the project. Keren Wen and Youjin Zhao designed the study and drafted the paper. Keren Wen, Youjin Zhao, Ziyu Zhu, Qian Li, Nanfang Pan and Shiqin Fu acquired and analyzed the data. Joaquim Radua, Eduard Vieta, Poomima Kumar, Graham J Kemp, Bharat B. Biswal, and Qiyong Gong critically revised the paper. Qiyong Gong gave final approval of the version to be published.

Conflict of interest statement

One of the authors, Dr Qiyong Gong, is also the editor-in-chief of Psychoradiology. He was blinded from reviewing or making decisions on the manuscript. Unrelated to the present work, Eduard Vieta has received grants and served as consultant, advisor or CME speaker for the following entities: Abbott, Allergan, Angelini, Dainippon Sumitomo Pharma, Janssen, Lundbeck, Novartis, Otsuka, Richter, Sage, Sanofi-Aventis, and Takeda. The other authors report no biomedical financial interests or potential conflicts of interest.

Acknowledgements

This study was supported by the National Natural Science Foundation of China (grant nos 81621003, 81761128023, 81820108018, 82027808, and 82001795) and NIH/NIMH R01MH112189-01. China Postdoctoral Science Foundation (2020M673245), Post-Doctoral Research Project of West China Hospital of Sichuan University (2021HXBH025), US-China joint grant (grant nos NSFC81761128023), Instituto de Salud Carlos III/European Union (ERDF/ESF, ‘Investing in your future’: CPI19/00009 and PII19/00394) and the project SLT006/17/00357, from PERIS 2016–2020 (Departament de Salut), CERCA Programme/Generalitat de Catalunya.

References

Allen P, Larsei F, McGuire PK, et al. (2008) The hallucinating brain: a review of structural and functional neuroimaging studies of hallucinations. Neurosci Biobehav Rev 32:175–91.
Ansell BR, Dwyer DB, Wood SJ, et al. (2015) Divergent effects of first-generation and second-generation antipsychotics on cortical thickness in first-episode psychosis. Psychol Med 45:515–27.
Bersani FS, Minichino A, Fojanesi M, et al. (2014) Cingulate cortex in schizophrenia: its relation with negative symptoms and psychotic onset. A review study. Eur Rev Med Pharmacol Sci 18:3354–67.
Bowd N, Hovington CL, Buchy L, et al. (2014) Cortical thinning in temporo-parietal junction (TPJ) in non-affective first-episode of psychosis patients with persistent negative symptoms. PLoS ONE 9:e101372.
Breitborde NJ, Srihari VH, Woods SW (2009) Review of the operational definition for first-episode psychosis. Early Interv Psychiatry 3:259–65.
Brent BK, Thermonos HW, Keshavan MS, et al. (2013) Gray matter alterations in schizophrenia high-risk youth and early-onset schizophrenia: a review of structural MRI findings. Child Adolesc Psychiatr Clin N Am 22:689–714.
Buchy L, Barbato M, Makowski C, et al. (2017) Mapping structural covariance networks of facial emotion recognition in early psychosis: a pilot study. Schizophr Res 189:146–52.
Buckner RL, Andrews-Hanna JR, Schacter DL (2008) The brain’s default network: anatomy, function, and relevance to disease. Ann N Y Acad Sci 1124:1–38.
Canal-Rivero M, Tordesillas-Gutiérrez D, Ruiz-Veguilla M, et al. (2020) Brain grey matter abnormalities in first episode non-affective psychosis patients with suicidal behaviours: the role of neurocognitive functioning. Prog Neuropsychopharmacol Biol Psychiatry 102:109948.
Chang JS, Choi S, Ha K, et al. (2011) Differential pattern of semantic memory organization between bipolar I and II disorders. Prog Neuropsychopharmacol Biol Psychiatry 35:1053–8.
Chudleigh C, Naismith SL, Blaszczynski A, et al. (2011) How does social functioning in the early stages of psychosis relate to depression and social anxiety? Early Interv Psychiatry 5:224–32.
Crow TJ, Chance SA, Priddle TH, et al. (2013) Laterality interacts with sex across the schizophrenia/bipolarity continuum: an interpretation of meta-analyses of structural MRI. Psychiatry Res 210:1234–44.
Cui Y, Liu B, Song M, et al. (2018) Auditory verbal hallucinations are related to cortical thinning in the left middle temporal gyrus of patients with schizophrenia. Psychol Med 48:115–22.
de Kovel CGF, Aftanas L, Aleman A, et al. (2019) No alterations of brain structural asymmetry in major depressive disorder: an ENIGMA Consortium Analysis. Am J Psychiatry 176:1039–49.
Downar J, Crawley AP, Mikulis DJ, et al. (2000) A multimodal cortical network for the detection of changes in the sensory environment. Nat Neurosci 3:277–83.
Dukart J, Smieszkova R, Harrisberger F, et al. (2017) Age-related brain structural alterations as an intermediate phenotype of psychosis. J Psychiatry Neurosci 42:307–19.
Eckert MA, Menon V, Walczak A, et al. (2009) At the heart of the ventral attention system: the right anterior insula. Hum Brain Mapp 30:2530–41.
Fischl B (2012) FreeSurfer. Neuroimage 62:774–81.
Fischl B, Dale AM (2000) Measuring the thickness of the human cerebral cortex from magnetic resonance images. Proc Natl Acad Sci USA 97:11050–5.

Fornito A, Yücel M, Dean B, et al. (2009) Anatomical abnormalities of the anterior cingulate cortex in schizophrenia: bridging the gap between neuroimaging and neuropathology. Schizophr Bull 35:973–93.

Fusar-Poli P, Smieskova R, Serafini G, et al. (2014) Neuroanatomical markers of genetic liability to psychosis and first episode psychosis: a voxelwise meta-analytical comparison. World J Biol Psychiatry 15:219–28.

Fusar-Poli P, Meyer-Lindenberg A (2013a) Striatal presynaptic dopamine in schizophrenia, part I: meta-analysis of dopamine active transporter (DAT) density. Schizophr Bull 39:22–32.

Fusar-Poli P, Meyer-Lindenberg A (2013b) Striatal presynaptic dopamine in schizophrenia, Part II: meta-analysis of [18F/11C]-DOPA PET studies. Schizophr Bull 39:33–42.

Ghasemi A, Zahediasl S (2012) Normality tests for statistical analysis: a guide for non-statisticians. Int J Endocrinol Metab 10:486–9.

Gong Q (2020) Psychoradiology. Neuroimaging Clinics of North America. 30:1–123. New York: Elsevier Inc.

Grant C, Addington J, Addington D, et al. (2001) Social functioning in first- and multiphase schizophrenia. Can J Psychiatry 46:746–9.

Hamilton LS, Narr KL, Luders E, et al. (2007) Asymmetries of cortical thickness: effects of handedness, sex, and schizophrenia. Neuroreport 18:1427–31.

Haring L, Muursepp A, Mottus R, et al. (2016) Cortical thickness and surface area correlates with cognitive dysfunction among first-episode psychosis patients. Psychol Med 46:2145–55.

Hui CL, Longenecker J, Wong GH, et al. (2012) Longitudinal changes in semantic categorization performance after symptomatic remission from first-episode psychosis: a 3-year follow-up study. Schizophr Res 137:118–23.

Hutton C, DeVita E, Ashburner J, et al. (2008) Voxel-based cortical thickness measurements in MRI. Neuroimage 40:1701–10.

Jang DP, Kim JJ, Chung TS, et al. (2006) Shape deformation of the insula in schizophrenia. Neuroimage 32:220–7.

Kay SR, Fiszbein A, Opler LA (1987) The positive and negative syndrome scale (PANSS) for schizophrenia. Schizophr Bull 13:261–76.

Konrad A, Winterer G (2008) Disturbed structural connectivity in schizophrenia primary factor in pathology or epiphenomenon? Schizophr Bull 34:72–92.

Kimmel CL, Alhassoon OM, Wollman SC, et al. (2016) Age-related parieto-occipital and other gray matter changes in borderline personality disorder: a meta-analysis of cortical and subcortical structures. Psychiatry Res Neuroimaging 251:15–25.

Konrad A, Winterer G (2008) Disturbed structural connectivity in schizophrenia primary factor in pathology or epiphenomenon? Schizophr Bull 34:72–92.

Leish TA, Tanase C, Geib BR, et al. (2015) A multimodal analysis of antipsychotic effects on brain structure and function in first-episode schizophrenia. JAMA Psychiatry 72:226–34.

Li F, Sun H, Biswal BB, et al. (2021) Artificial intelligence applications in psychoradiology. Psychoradiology 1:94–107.

Li Q, Zhao Y, Chen Z, et al. (2020) Meta-analysis of cortical thickness abnormalities in medication-free patients with major depressive disorder. Neuropsychopharmacology 45:703–12.

Lin Y, Li M, Zhou Y, et al. (2019) Age-related reduction in cortical thickness in first-episode treatment-naïve patients with schizophrenia. Neurosci Bull 35:688–96.

Lui S, Zhou XJ, Sweeney JA, et al. (2016) Psychoradiology: the frontier of neuroimaging in psychiatry. Radiology 281:357–72.

Malikarjun PK, Lalousis PA, Dunne TF, et al. (2018) Aberrant salience network functional connectivity in auditory verbal hallucinations: a first episode psychosis sample. Transl Psychiatry 8:69.

Menon V, Uddin LQ (2010) Saliency, switching, attention and control: a network model of insula function. Brain Struct Funct 214:655–67.

Merritt K, Luque Laguna P, Irfan A, et al. (2021) Longitudinal structural MRI findings in individuals at genetic and clinical high risk for psychosis: a systematic review. Front Psychiatry 12:620401.

Mitchell RJ, Crow TJ (2005) Right hemisphere language functions and schizophrenia: the forgotten hemisphere? Brain 128:963–78.

Morosini P, Magliano L, Brambilla L, et al. (2000) Development, reliability and acceptability of a new version of the DSM-IV Social and Occupational Functioning Assessment Scale (SOFAS) to assess routine social functioning. Acta Psychiatr Scand 101:323–9.

Mulders PC, van Eijndhoven PF, Schene AH, et al. (2015) Resting-state functional connectivity in major depressive disorder: a review. Neurosci Biobehav Rev 56:330–44.

Nasrallah HA (1985) The unintegrated right cerebral hemisphere: a possible mechanism for Schneiderian delusions in schizophrenia. Compr Psychiatry 26:273–82.

O’Neill A, Mechelli A, Bhattacharyya S (2019) Disconnection of large-scale functional networks in early psychosis: a meta-analysis. Schizophr Bull 45:579–90.

Palaniyappan L, Liddle PF (2012a) Do the salience network play a cardinal role in psychosis? An emerging hypothesis of insular dysfunction. J Psychiatry Neurosci 37:399–406.

Palaniyappan L, Liddle PF (2012b) Does the salience network play a cardinal role in psychosis? An emerging hypothesis of insular dysfunction. J Psychiatry Neurosci 37:399–406.

Poppa T, Bechara A (2018) The somatic marker hypothesis: revisiting the role of the ‘body-loop’ in decision-making. Curr Opin Behav Sci 19:61–6.

Quah SR, Cockerham WC (eds.) (2017) International Encyclopedia of Public Health, 2nd edn: Academic Press.

Radua J, Borgwardt S, Crescini A, et al. (2012a) Multimodal meta-analysis of structural and functional brain changes in first...
episode psychosis and the effects of antipsychotic mediation. *Neurosci Biobehav Rev* 36:2325–33.

Radua J, Del Pozo NO, Gómez J, et al. (2014a) Meta-analysis of functional neuroimaging studies indicates that an increase of cognitive difficulty during executive tasks engages brain regions associated with time perception. *Neuropsychologia* 58:14–22.

Radua J, Mataix-Cols D, Phillips ML, et al. (2012b) A new meta-analytic method for neuroimaging studies that combines reported peak coordinates and statistical parametric maps. *Eur Psychiatry* 27:605–11.

Radua J, Via E, Catani M, et al. (2011) Voxel-based meta-analysis of regional white-matter volume differences in autism spectrum disorder versus healthy controls. *Psychol Med* 41:1539–50.

Radua J, Mataix-Cols D (2009) Voxel-wise meta-analysis of grey matter changes in obsessive–compulsive disorder. *Br J Psychiatry* 195:393–402.

Radua J, Rubia K, Canales-Rodríguez EJ, et al. (2014b) Anisotropic kernels for coordinate-based meta-analyses of neuroimaging studies. *Front Psychiatry* 5:13.

Rais M, van Haren NEM, Cahn W, et al. (2010) Cannabis use and progressive cortical thickness loss in areas rich in CB1 receptors during the first five years of schizophrenia. *Eur Neuropsychopharmacol* 20:855–65.

Reed SI (2008) First-episode psychosis: a literature review. *Int J Ment Health Nurs* 17:85–91.

Reniers RLEP, Garner B, Phassouliotis C, et al. (2015) The relationship between stress, HPA axis functioning and brain structure in first episode psychosis over the first 12 weeks of treatment. *Psychiatry Res Neuroimaging* 231:111–9.

Roiz-Santíanez R, Perez-Iglesias R, Quintero C, et al. (2010) Insular cortex thinning in first episode schizophrenia patients. *Psychiatry Res Neuroimaging* 182:216–22.

Rossell SL, Rabe-Hesketh SS, Shapleske JS, et al. (1999) Is semantic fluency differentially impaired in schizophrenic patients with delusions? *J Clin Exp Neuropsychol* 21:629–42.

Scanlon C, Anderson-Schmidt H, Kilmarin L, et al. (2014) Cortical thinning and caudate abnormalities in first episode psychosis and their association with clinical outcome. *Schizophr Res* 159:36–42.

Schober P, Boer C, Schwarte LA (2018) Correlation coefficients: appropriate use and interpretation. *Anesth Analg* 126:1763–8.

Schultz CC, Koch K, Wagner G, et al. (2010) Reduced cortical thickness in first episode schizophrenia. *Schizophr Res* 116:204–9.

Shah C, Zhang W, Xiao Y, et al. (2017) Common pattern of gray-matter abnormalities in drug-naive and medicated first-episode schizophrenia: a multimodal meta-analysis. *Psychol Med* 47:401–13.

Simon GE, Stewart C, Yarborough Bjo, et al. (2018) Mortality rates after the first diagnosis of psychotic disorder in adolescents and young adults. *JAMA Psychiatry* 75:254–60.

Song X, Quan M, Lv L, et al. (2015) Decreased cortical thickness in drug naive first episode schizophrenia: in relation to serum levels of BDNF. *J Psychiatr Res* 60:22–8.

Sowell ER, Thompson PM, Leonard CM, et al. (2004) Longitudinal mapping of cortical thickness and brain growth in normal children. *J Neurosci* 24:8223–31.

Squire LR (1992) Declarative and nondeclarative memory: multiple brain systems supporting learning and memory. *J Cogn Neurosci* 4:232–43.

Sun H, Lui S, Yao L, et al. (2015) Two patterns of white matter abnormalities in medication-naive patients with first-episode schizophrenia revealed by diffusion tensor imaging and cluster analysis. *JAMA Psychiatry* 72:678–86.

Sun J, Maller JJ, Guo L, et al. (2009) Superior temporal gyrus volume change in schizophrenia: a review on region of interest volumetric studies. *Brain Res Rev* 61:14–32.

Tan EJ, Rossell SL (2014) Building a neurocognitive profile of thought disorder in schizophrenia using a standardized test battery. *Schizophr Res* 152:242–5.

Tan EJ, Neill E, Tomlinson K, et al. (2020) Semantic memory impairment across the schizophrenia continuum: a meta-analysis of category fluency performance. *Schizophr Bull* 1.

Todtenkopf MS, Vincent SL, Benes FM (2005) A cross-study meta-analysis and three-dimensional comparison of cell counting in the anterior cingulate cortex of schizophrenic and bipolar brain. *Schizophr Res* 73:79–89.

Tordesillas-Gutierrez D, Ayasa-Arriola R, Delgado-Alvarado M, et al. (2018) The right occipital lobe and poor insight in first episode psychosis. *PLoS ONE* 13:e0197715.

Vieira S, Gong QY, Pinaya WHL, et al. (2020) Using machine learning and structural neuroimaging to detect first episode psychosis: reconsidering the evidence. *Schizophr Bull* 46:17–26.

Walton E, Hibar DP, van Erp TG, et al. (2017) Positive symptoms associate with cortical thinning in the superior temporal gyrus via the ENIGMA Schizophrenia consortium. *Acta Psychiatri Scand* 135:439–47.

Wannan CMJ, Cropley VL, Chakravarty MM, et al. (2019) Evidence for network-based cortical thickness reductions in schizophrenia. *Am J Psychiatry* 176:552–63.

Wells GA, Shea B, O’Connell D, et al. (2000) The Newcastle-Ottawa Scale (NOS) for assessing the quality of nonrandomised studies in meta-analyses.

Winkler AM, Kochunov P, Blangero J, et al. (2010) Cortical thickness or grey matter volume? The importance of selecting the phenotype for imaging genetics studies. *Neuroimage* 53:1135–46.

Wylie KP, Tregellas JR (2010) The role of the insula in schizophrenia. *Schizophr Res* 123:93–104.

Xiao Y, Lui S, Deng W, et al. (2015) Altered cortical thickness related to clinical severity but not the untreated disease duration in schizophrenia. *Schizophr Bull* 41:201–10.

Xu J, Wang J, Fan L, et al. (2015) Tractography-based parcelation of the human middle temporal gyrus. *Sci Rep* 5:18883.

Zhang H, Li L, Wu M, et al. (2016) Brain gray matter alterations in first episodes of depression: a meta-analysis of whole-brain studies. *Neurosci Biobehav Rev* 60:43–50.