Research article

Large-scale networks underlie cognitive insight differs between untreated adolescents ongoing their first schizophrenic episode and their reference non-schizophrenic mates

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ABSTRACT

Objectives: Cognitive insight (CI), the ability to perceive erroneous beliefs and correcting them based on safe experiences, is a common cognitive manifestation among schizophrenic individuals. Even though the functional morphology of the default mode network (DMN), the central executive network (CEN) and the salience network (SN) differs between non-schizophrenic and schizophrenic individuals, it is unclear whether such differences are already in place by the first schizophrenic episode.

Methods: Forty-two adolescents, including twenty-one AOS subjects was recruited, and performed independent component analysis (ICA) on resting-state fMRI data to explore alterations in the three networks in schizophrenia and the association of network changes with Beck Cognitive Insight Scale (BCIS) scores.

Results: Compared to the non-schizophrenic group, the AOS group showed hyper-connectivity in the left middle temporal gyrus (MTG) and hypo-connectivity in the right parahippocampal gyrus within the DMN; hypo-connectivity in the dorsal anterior cingulate (dACC) and supplementary motor area (SMA) within the SN were also detected in AOS individuals. CI subscores were positively correlated with functional connectivity (FC) in the right parahippocampal gyrus.

Conclusions: The correlations reported here suggest that increased DMN connectivity in the right parahippocampal gyrus might be an early neural correlate of reduced cognitive insight in a number, but not all, adolescent untreated individuals ongoing their first schizophrenic episode.

1. Introduction

The term schizophrenia encompasses a set of phenotypes whose lifetime prevalence reaches up to 0.6% in China (Huang et al., 2019), often characterized as dysfunction of cognitive, emotion and behaviour. Commonly, individuals ongoing their first schizophrenic episode during adolescence (AOS) display such event between 13 and 18 years of age. AOS individuals may exhibit genetic markers, disruptive social relationships, and poor responses to antipsychotics and psychotherapy. Lack of insight is a core manifestation of schizophrenia and indicates that individuals do not have a complete awareness of their psychotic disorder (Pick, 1882). With in-depth study, insight is more likely to be regarded as a neurocognition-involved issue. Beck and his colleagues proposed the concept of cognitive insight (CI), coupled with an additional cognitive dimension, defined as the ability to detect and correct one’s erroneous beliefs, and developed the Beck Cognitive Insight Scale (BCIS) to assess the capacity for reevaluating anomalous experiences and misinterpretations (Beck et al., 2004). Compared with clinical insight,
cognitive insight tends to be more stable and less impacted by the duration of illness, number of hospitalizations, age and years of education (Bora et al., 2007; Kim et al., 2015a).

Schizophrenia is thought to be the result of brain-wide functional dysconnectivity, and this hypothesis is supported by various neuroimaging evidence. Resting-state fMRI offers an approach to capture intrinsic spatiotemporal brain connectivity dynamics, which can reflect fundamental functional characteristics of the brain. The well-recognized resting-state networks mainly consist of the default mode network (DMN), executive control network (ECN), salience network (SN), dorsal attention network (DAN), and self-referential network (SRN). Dysfunction in one core network is often coupled with abnormalities in the other networks (Menon, 2011). The anomalous integration of information within and across these networks is important in linking the associated impaired cognitive features characteristic of psychopathology (Buckholtz and Meyer-Lindenberg, 2012; Gong et al., 2017). What is noteworthy is that aberrant resting state networks (RSNs) have a close relationship with self-related information processes (Camchong et al., 2011; Liu et al., 2012). Previous studies have claimed that cognitive insight is highly related to self-reflection, so we speculated that RSN dysfunction may cause cognitive insight disruption.

To further the understanding of how RSN function is relevant to phenotype with schizophrenia, the functional model was characterized in a development cohort. During this critical node of brain maturation progress, developmental delays get to crucial milestones, progressive grey matter volume loss and aberrant functional connectivity have been detected in early-onset case (Jacobsen and Rapoport, 1998; Tang et al., 2013), suggest a potentially deficit in information integrate and cognitive progress. Moreover, compared with adult-onset schizophrenia, early-onset schizophrenia appears to be more insidious (Kolvin et al., 1971; Eggers, 1978), often combine with premorbid maladjustment. The undiscovered cognitive impairment before illness onset might basis for decision making and self-judgment. Above all, we hypothesized a correlation between the alternative functional mode and the lack of cognitive insight in AOS.

Thus, in the present study, the authors aimed to explore the differences in network patterns in AOS individuals by using independent component analysis (ICA) with a main focus on the triple-network model. Furthermore, correlation analyses were applied to explore whether this is related to CI in AOS. It was hypothesized that AOS subjects may differ from non-schizophrenic in FC in the triple-network model, and those alterations may contribute to impaired CI.

### 2.2. Data acquisition

All participants underwent a resting-state echo-planar imaging (EPI) scan through a 3.0T Siemens MRI scanner. They were informed to close their eyes, relax, stay awake, and avoid thinking and moving. Their heads were fixed by foam carpet to reduce head movement during the scan, and sound-insulated earphones were worn to prevent noise interference. Functional images were acquired with the following parameters: time points = 255, repetition time (TR) = 2000 ms, echo time (TE) = 30 m s, slice thickness = 4 mm, 35 axial slices, matrix:64 mm × 64 mm, flip angle: 90°, and field of view (FOV) = 240 mm × 240 mm. High-resolution magnetization-prepared fast gradient echo (MP-FGRE) T1-weighted images were also acquired (TR = 1900 m s, TE = 2.25 m s, slice thickness = 1 mm, matrix: 256 mm × 256 mm, and flip angle: 9°).

### 2.3. Image pre-processing

The functional images were preprocessed using the Data Processing Assistant for Resting-State fMRI (DPARSF version 4.4), which synthesizes procedures in statistical parametric mapping (SPM12, http://www.fil.ion.ucl.ac.uk/spm/software/spm12). The first 10 timepoints of the functional images were removed to ensure that all the images were acquired in a stable condition. The remaining 245 time points were corrected for slice-time differences and realigned to the first image. Structural images were co-registered to the mean functional image. The transformed structural images were then segmented into gray matter, white matter and cerebrospinal fluid (CSF). Then, mean time signals were extracted and regressed out. A higher-level Friston-24 model was used to regress head motion effects out of the realigned data (the 24 parameters included 6 head motion parameters, 6 head motion parameters one time point before, and the 12 corresponding squared items). The mean framewise displacement (FD) was further calculated as a measure of the microscale head motion of each subject. Data from subjects with high head movement (FD > 0.2 mm) were excluded. All images were spatially normalized into Montreal Neurological Institute (MNI) space (voxel size = 3 mm × 3 mm ×3 mm). Finally, the images were smoothed with an 8-mm full-width half-maximum (FWHM) Gaussian kernel.

### 2.4. Independent component analysis

The ICA method was used for the preprocessed data with the Group ICA FMRI Toolbox (GIFT; http://icatb.sourceforge.net/gift/gift_start up.php). The individual components were estimated using the maximum description length (MDL) criterion. Data reduction was performed with a two-stage principal component analysis (PCA) that consisted of individual-level analysis and group-level analysis. After reduction, the infomax algorithm was used to decompose the data into a fixed set of 20 spatially independent components, and then the default method in the GIFT toolbox was used to test the stability of those components. Because the components we obtained from the above step belong to the group but not individuals, the data were

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back-reconstructed to obtain the spatial maps and time courses of individual components. The independent components were subsequently categorized into different intrinsic functional networks based on spatial correlation ($r > 0.2$) with a priori templates (Beckmann et al., 2005; Smith et al., 2009), and then visually confirmed the results. Finally, three intrinsic functional networks was identified with 6 independent components.

### 2.5. Statistical analyses

Spatial maps of selected components were compared between subjects and non-schizophrenic separately to acquire the components. Then, the common parts of the masks from the AOS group and non-schizophrenic group were used to create one mask for further group comparison.

To find the differences in spatial maps between the AOS and non-schizophrenic groups, components from both groups were entered in a two-sample t-test with the previously constructed mask. Next, the statistical significance of the results was defined using a threshold of $p < 0.01$ (uncorrected) at the voxel level and AlphaSim-corrected $p < 0.05$ at the cluster level. A functional network connectivity (FNC) correlation matrix was created to investigate FC between these three networks using the MANCOVA toolbox. Two-sample t-tests were applied to compare FNC correlations between the AOS and non-schizophrenic groups using false discovery rate (FDR) ($q = 0.05$) correction.

Demographic data, such as age, sex and years of education, were compared using t-tests or chi-square tests. Pearson correlation analyses were used between CI and clinical data, and FDR adjustment was applied ($p < 0.05$). Linear correlation analyses were used to explore the relationships between the FC of target brain areas and the clinical symptoms and CI.

### 3. Results

#### 3.1. Demographics and clinical characteristics

The AOS group and non-schizophrenic group did not have significant difference in age ($t = -1.360, p = 0.182$), sex ratio ($\chi^2 = 4.29, p = 0.041$), years of education ($t = -1.21, p = 0.234$), BCIS Self-reactiveness score ($t = 0.345, p = 0.732$), Self-certainty score ($t = -2.116, p = 0.041$), Composite Index ($t = 1.954, p = 0.058$), SAPS scores ($t = 0.345, p = 0.732$), SANS scores ($t = 0.345, p = 0.732$), and PSYRAT scores ($t = 0.345, p = 0.732$). However, the AOS group showed a significant increase in B) SC scores. No significant difference was found in A) SR scores or C) composite index.

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### Table 1. Clinical and demographic characteristics of AOS individuals and non-schizophrenic.

|                      | HC group | AOS group | $t$/ $\chi^2$ | $p$ |
|----------------------|----------|-----------|---------------|-----|
| Age (years)          | 16.4 ± 1.0 | 16.9 ± 1.3 | -1.360        | 0.182 |
| Sex (male/female)    | 8/13     | 6/15      | 0.429         | 0.513 |
| Years of education (years) | 10.1 ± 1.3 | 10.6 ± 1.5 | -1.21        | 0.234 |
| BCIS                 |          |           |               |      |
| Self-reactiveness (SR) | 15.1 ± 3.2 | 14.8 ± 3.9 | 0.345         | 0.732 |
| Self-certainty (SC)  | 7.3 ± 2.4 | 9.2 ± 3.4  | -2.116        | 0.041 |
| Composite Index      | 7.8 ± 4.1 | 5.5 ± 3.4  | 1.954         | 0.058 |
| SAPS                 |          | 28.38 ± 13.02 |          |      |
| SANS                 |          | 33.57 ± 25.65 |          |      |
| PSYRAT               |          | 18.10 ± 16.91 |          |      |

SR, self-reactiveness; SC, self-certainty; SAPS, Scale for Assessment of Positive Symptoms; SANS: Scale for Assessment of Negative Symptoms; PSYRAT, Psychotic Rating Scales.

* AOS > Controls ($p < 0.05$).
0.513), or years of education ($t = -1.210, p = 0.234$). The mean SAPS, SANS and BCIS scores of these two groups are shown in Table 1. The AOS group showed significantly higher SC scores than the non-schizophrenic group ($t = -2.116, p = 0.041$) (Figure 1B), there was no significant difference in SR scores and composite index (Figure 1A, 1C). The Pearson correlation analysis between cognitive insight and clinical scale values showed that there were no significant result at the FDR corrected thresholds (Table 2).

### 3.2. Within-network dysconnectivity

The intrinsic functional networks extracted by group ICA are presented in Figure 2 Compared to the non-schizophrenic group, the AOS group showed imbalanced FC within the DMN with weaker connectivity at the level of the left middle temporal gyrus (MTG) but stronger connectivity for the right parahippocampal gyrus. In addition, the AOS group had weaker FC within the SN in the dorsal anterior cingulate cortex (dACC) and supplementary motor area (SMA) (voxel-level $p < 0.01$, cluster-level $p < 0.05$, AlphaSim correction) (Figure 3, Table 3). No significant results were found within the CEN between these two groups. Further analysis showed that the SC scores were positively related to the FC strength of the right parahippocampal gyrus ($r = 0.451, p < 0.05$) (Figure 4).

### 3.3. Between-network dysconnectivity

There were no significant differences in between-network connectivity between the AOS and non-schizophrenic groups.

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**Table 2. Correlation analysis* between cognitive insight and clinical scales.**

|                  | SR        | SC        | composite index |
|------------------|-----------|-----------|-----------------|
| Age (years) (P)  | 0.322 (0.385) | 0.274 (0.491) | 0.099 (0.675) |
| Years of education (years) (P) | 0.417 (0.300) | 0.343 (0.385) | 0.139 (0.631) |
| SAPS (P)         | -0.249 (0.519) | -0.097 (0.675) | -0.190 (0.558) |
| SANS (P)         | -0.192 (0.558) | -0.471 (0.300) | 0.249 (0.519) |
| PSYRAT (P)       | 0.423 (0.300) | 0.162 (0.604) | 0.327 (0.385) |

SR, self-reflectiveness; SC, self-certainty; SAPS, Scale for Assessment of Positive Symptoms; SANS, Scale for Assessment of Negative Symptoms; PSYRAT, Psychotic Rating Scales. * at the FDR correction threshold.

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![Figure 2. Three intrinsic functional networks extracted by group independent component analysis (ICA). Abbreviations: DMN, default mode network; CEN, central executive network; SN, salience network.](image-url)
4. Discussion

Schizophrenia is defined as a kind of severe mental disorder characterized by deficit in the relation between thought, emotion, and behaviour, often accompany with impaired cognitive insight. As expected, AOS demonstrated alternative functional connectivity in intrinsic neural network, and this differs were associated with cognitive insight. This results indicated that the alternative functioning of the right parahippocampal gyrus within the DMN probably lead to improper self-referential information process, might be the potential foundation of CI impairments. To our knowledge, no previous work have evaluated whether reduced cognitive insight is already present in adolescent untreated individuals ongoing their first schizophrenic episode.

The only study exploring the relationship between CI and RSN found that CI was associated with functional connectivity in the right inferior frontal cortex (IFC) and left ACC in adult schizophrenia subjects (Gerritsen et al., 2014). This study presented hyper-connectivity in the right parahippocampal gyrus within the DMN in AOS individuals, that increased connectivity in the AOS subjects compared to the non-schizophrenic was related to higher SC scores. Authors have suggested that the parahippocampal gyrus is engaged extensively in cognitive processing such as memory recollection (Gary and Van, 1982; Eichenbaum et al., 1996), hallucinations (Diederen et al., 2010), and emotion (Arias et al., 2015; Kim et al., 2015b). Besides, the medial temporal subsystem in DMN, including the parahippocampal gyrus, has
been implicated in past and future autobiographical thought and contextual retrieval (Andrews-Hanna et al., 2014). One study with self-judgment reported co-activation between the VMPFC and parahippocampus gyrus while making self-judgments, prompting the retrieval of specific autobiographical details to influence self-referential information assessments while making self-judgments (Feyers et al., 2010). This research was predicated on the theory that RSNs dysfunction contribute to deficit self-related information processes. According to the description, the SC score refers to self-evaluation and self-judgment, which fall largely in the domain of self-reference. As mentioned before, parahippocampal gyrus plays a critical role in autobiographical memory and contextual information processing, it can be inferred that the parahippocampal gyrus participates in self-judgment through its engagement in autobiographical memory, which could explain the positive correlation that found between SC scores and FC in the parahippocampal gyrus. This work employed first-onset drug-naive AOS individuals and provides evidence that increased DMN connectivity in the right parahippocampal gyrus might be an indication of the reduction of cognitive insight in individuals during development.

The DMN is well known for its involvement in self-referential (Gusnard et al., 2001), integrative processes (Greicius et al., 2003), internal emotion (Raichle and Snyder, 2007) and internal thought (Mason et al., 2007). In the current study, the dysconnectivity in the left MTG within DMN are consistent with previous resting-state studies with AOS subjects (Zhang et al., 2017). As an important region of the DMN, abundant evidences has emphasized that the MTG is involved in important cognitive processes (Cabeza and Nyberg, 2000; Kristinsson et al., 2020) and plays an influential role in the activation of psychiatric symptoms (Zhang et al., 2017; Cui et al., 2018). Otherwise, the MTG has been suggested to be crucial in self-representations by retrieval information from autobiographical memory (Feng et al., 2018). On the other hand, AOS subjects presented significant hypo-connectivity in the dACC and SMA within the SN. These results are in line with previous studies with schizophrenia (Pu et al., 2012), while no relevant study with AOS subject can be identified. The SN detects the salience of internal and external stimuli (Sridharan et al., 2008; Menon and Uddin, 2010), alternative attribution of salience to external and internal stimuli is thought to arouse psychotic symptoms. For instance, Malikkarjun et al. (2018) reported dysconnectivity between the SN and DMN in first-episode schizophrenia with auditory verbal hallucinations, similar results also appeared in chronic schizophrenia (Alonso-Solis et al., 2015) and individuals with PTSD (Cheng et al., 2019). Moreover, variety in interactions between the SN and prefrontal system, which may lead to an unbalanced salience-execution loop, have been hypothesized to underlie the neural background of psychosis (Palaniyappan et al., 2013). In addition, alternative integration between the SN and DMN and the CEN in schizophrenia have already reported (Wu and Jiang, 2019; Manoliu et al., 2014), implied that the ability to coordinate self-monitoring and task performance was impaired in schizophrenia individuals; thus, the reduced requirement of the ACC and SMA suggest a disturbance in this system.

This research failed to find a significant difference in the CEN between AOS individuals and non-schizophrenic. The CEN is mainly involved in working memory and decision making (Michael, 2005; Koechlin and Summerfield, 2007), dysfunction in the CEN has been widely reported in schizophrenia, and both hyper-connectivity (Krishnadass et al., 2014) and hypo-connectivity (Woodward et al., 2011) have been found. However, parallel with the development of the neural system, brain networks are also altered during adolescence (Menon, 2013). Thus, AOS allows researchers to treat schizophrenia as alternative phenotypes of brain connectivity emerging from divergent neurodevelopmental trajectories. Based on the small sample size, it is hard to support the hypothesis of CEN dysfunction in AOS, amplifying the quantity of subjects may have a different result.

Although triple-network model alterations was examined on a large scale with reliable measurements in an adolescent population with schizophrenia, the present study still had several limitations. First, restricting the sample to drug-naive adolescents enhanced the homogeneity but limited the generalizability of our results to adult individuals. Second, the sample size was relatively small, and future fMRI studies in a larger sample will be needed to validate these findings. Last, the conclusion is based on adolescent-onset untreated individuals ongoing their first schizophrenic episode, for the reason that schizophrenia might be a continuum of phenotypes, only in a certain percentage increased DMN connectivity in the right parahippocampal gyrus could be treated as early neural marker of impaired cognitive insight.

In conclusion, we characterized the large-scale network changes underlying CI in untreated adolescents ongoing their first schizophrenic episode. In line with previous predictions, this results showed that dysfunction in the right parahippocampal gyrus within the DMN might be an early neural correlate of reduced cognitive insight, provide neurobiological evidence to support the theory of brain-behaviors relationships is critical for cognitive process and clinical symptoms, offer fundamental information for further research.

Declarations

Author contribution statement

Guoping Huang; Xiaoxi Huang: Conceived and designed the experiments; Wrote the paper.
Ruofei Ji; Ming Zhou: Performed the experiments; Analyzed and interpreted the data; Wrote the paper.
Na Ou; Hudan Chen; Yang Li: Contributed reagents, materials, analysis tools or data; Wrote the paper.

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Data availability statement

The authors do not have permission to share data.

Declaration of interest’s statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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