


Altered brain structural and functional connectivity in schizotypy

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Original Article

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Abstract

Background. Schizotypy refers to schizophrenia-like traits below the clinical threshold in the general population. The pathological development of schizophrenia has been postulated to evolve from the initial coexistence of ‘brain disconnection’ and ‘brain connectivity compensation’ to ‘brain connectivity decompensation’.

Methods. In this study, we examined the brain connectivity changes associated with schizotypy by combining brain white matter structural connectivity, static and dynamic functional connectivity analysis of diffusion tensor imaging data and resting-state functional magnetic resonance imaging data. A total of 87 participants with a high level of schizotypal traits and 122 control participants completed the experiment. Group differences in whole-brain white matter structural connectivity probability, static mean functional connectivity strength, dynamic functional connectivity variability and stability among 264 brain sub-regions of interests were investigated.

Results. We found that individuals with high schizotypy exhibited increased structural connectivity probability within the task control network and within the default mode network; increased variability and decreased stability of functional connectivity within the default mode network and between the auditory network and the subcortical network; and decreased static mean functional connectivity strength mainly associated with the sensorimotor network, the default mode network and the task control network.

Conclusions. These findings highlight the specific changes in brain connectivity associated with schizotypy and indicate that both decompensatory and compensatory changes in structural connectivity within the default mode network and the task control network in the context of whole-brain functional disconnection may be an important neurobiological correlate in individuals with high schizotypy.

Introduction

Schizotypy refers to subclinical schizophrenia-like traits found in the general population (Meehl, 1962; Sarig, Dar, & Liberman, 2012). Its clinical manifestations include cognitive-perceptual (positive), interpersonal (negative), and cognitive disorganized features (Polner *et al.*, 2019; Raine *et al.*, 1994), which are qualitatively similar to schizophrenia symptoms and correlate with schizophrenia liability (Grant, Green, & Mason, 2018). Schizotypy may represent an ‘endophenotype’ on the path to schizophrenia (Grant, 2015), and can be quantitatively assessed using the Schizotypal Personality Questionnaire (SPQ; Raine, 1991). Epidemiological studies have shown that the proportion of individuals with a high level of schizotypal traits in the general population is about 10% (Ettinger, Meyhöfer, Steffens, Wagner, & Koutsouleris, 2014; Lenzenweger, 2011). Studying schizotypy could help in understanding schizophrenia-related psychopathology (Kwapil & Barrantes-Vidal, 2014) and putative schizophrenia endophenotypes (Lenzenweger, 2015).

Previous studies have suggested that the neurobiological basis of schizophrenia may be ‘brain disconnection’ or ‘brain dysconnection’ (Friston, Brown, Siemerikus, & Stephan, 2016; Friston & Frith, 1995; Stephan, Friston, & Frith, 2009; Wang *et al.*, 2018b). Specifically, one study has found that fractional anisotropy values of the fronto-temporal white matter tracts correlated negatively with scores on the SPQ cognitive-perceptual subscale in the general population (Nelson *et al.*, 2011). Resting-state functional magnetic resonance imaging (rs-fMRI) studies have found that individuals with a high level of schizotypal traits exhibited reduced functional connectivity between the precuneus and the parahippocampus, which was

negatively correlated with SPQ total scores (Zhu et al., 2017). In the general population, SPQ cognitive-perceptual subscale scores correlated positively with functional connectivity between the ventral striatum and the frontal cortex, and negatively with the functional connectivity between the dorsal striatum and the posterior cingulate cortex (Wang, Ettinger, Meindl, & Chan, 2018a, 2018b). An independent component analysis study found that SPQ cognitive-perceptual subscale scores correlated positively with low-frequency oscillations of the visual network, while SPQ interpersonal subscale scores correlated negatively with low-frequency oscillations of the auditory network in the general population (Lagioia, Van De Ville, Debbané, Lazeyras, & Eliez, 2010). These findings suggest that the development of schizotypy may be related to changed brain network connectivity (Mohr & Claridge, 2015). Comprehensive investigation in individuals with a high level of schizotypal traits could enhance our understanding of the differences and similarities in structural and functional connectivity changes associated with schizotypy and could identify which of these changes are important in individuals with high schizotypy (Lenzenweger, 2006).

Previous studies on schizotypy have also paid little attention to the time dynamics of functional connectivity. Temporal coupling between brain sub-regions is time-variant rather than time-invariant (Nomi et al., 2017). Dynamic functional connectivity (DFC), also called 'chronnectome', could provide insight into the mutability of interacted functional connectivity networks. One previous study has found that schizophrenia patients exhibited dynamic changes in thalamic hyperconnectivity and sensory hypoconnectivity (Damaraju et al., 2014). Another study found that individuals with psychotic-like experiences exhibited altered brain dynamics associated with hyperconnectivity of the visual network and hypoconnectivity of the default mode network (Barber, Lindquist, DeRosse, & Karlsgodt, 2018). Overall, the stability of DFC in individuals with high level of schizotypal traits warrants further investigation.

In this study, we investigated the structural connectivity, static functional connectivity and DFC using diffusion tensor imaging (DTI), and resting-state functional magnetic resonance imaging (rs-fMRI) technology. We estimated structural connectivity by probabilistic tractography and DFC by a new and reliable Dynamic Conditional Correlation (DCC) method (Lindquist, Xu, Nebel, & Caffo, 2014), which is more reliable than slide-window and non-parametric estimation methods with a quantitative description of time-varying characteristics (Choe et al., 2017). The representative whole-brain template we used in this study was identified by Power et al. (2011), which included 264 regions of interest (ROI) belonging to diverse brain networks. The structural connectivity probability, the static mean functional connectivity strength and DFC indices including the DFC variability and the DFC stability between these ROIs were calculated.

Based on previous findings of both deficits and compensatory brain connectivity changes between brain regions or networks associated with the development of schizophrenia spectrum disorder (Abram et al., 2017; Andreasen, Paradiso, & O'leary, 1998; Chan et al., 2011; Cho et al., 2015; Pettersson-Yeo, Allen, Benetti, McGuire, & Mechelli, 2011), we propose a neurobiological developmental possibility of schizophrenia called the 'Connectivity Decompensation Hypothesis', which hypothesize that the pathological development of schizophrenia spectrum continuum is a gradual process from functional compensation to partial decompensation to complete decompensation. In this model, some neural circuits associated with emotion, sensory

and perceptual inference may gradually become impaired in susceptible individuals due to a combination of genetic and/or undesirable internal and external environmental factors (Friston et al., 2016). These abnormalities could still be partially compensated initially and these individuals could maintain normal functioning by gradual structural proliferation, remyelination, adaptive functional adjustment or reorganization of brain sub-regions associated with cognitive regulation and self-monitoring (Mohr & Claridge, 2015). The majority of them would never develop the illness. However, if these predisposing factors are excessive and persistent, or if neural compensation mechanisms fail, psychotic-like features and cognitive function maladjustment could be further aggravated, leading to clinical illness onset. Individuals with a high level of schizotypal traits are in the early stage of this process, and investigation of them could offer the best opportunity for early intervention and prevention and avoid confounding factors such as antipsychotic medication exposure and psychiatric comorbidities (Barrantes-Vidal, Grant, & Kwapil, 2015).

Thus, the objectives of our study were (1) to investigate common and specific changes in functional and structural connectivity in individuals with a high level of schizotypal traits compared with controls; and (2) to verify the connectivity decompensation hypothesis in individuals with a high level of schizotypal traits. If functional and structural connectivity between ROIs were both increased in the high schizotypy group, it could be considered as a compensation effect in this pathway. If functional connectivity between ROIs were decreased while structural connectivity was increased, it could imply decompensation. Based on previous studies, we hypothesized that changes in structural and functional connectivity in individuals with a high level of schizotypal traits would exhibit both decompensatory and compensatory effects, which could indicate 'brain disconnection'.

Material and methods

Participants

We first collected 4000 online schizotypal personality questionnaire (SPQ) responses from various universities randomly in Beijing through advertisement in 2018. Every voluntary participant was a college student and was paid. The 90th percentile of the SPQ total score (41 points) among these participants was taken as the cut-off score, which was in accord with previous studies setting individuals scoring in the top 10% of SPQ scores in the general population as the high schizotypy group (Chan et al., 2011; Ettinger et al., 2014; Lenzenweger, 2011; Shan et al., 2020; Shi et al., 2017; Wang et al., 2014, 2020a, 2020b). Students with a score above 41 were assigned to the high schizotypy group, and those with a score below 41 were assigned as controls. Then individuals allocated to the high schizotypy group and a corresponding number of controls were contacted by text message to invite them to participate in the subsequent MRI experiment. Of the total, 90% of the participants responded to the online survey. Of which, 246 responders agreed to undertake the brain scan. Inclusion criteria for all participants included Han ethnicity and aged 18 years or above. Exclusion criteria for all participants included any diagnosis of physical or psychiatric disorders and substance dependence, any family history of psychiatric disorder and any contraindications for MRI scanning. In addition, any participant who did not complete all study procedures, whose imaging quality was poor, and who had head motions greater

than 2 mm and/or 2° during scanning were excluded. With these criteria, 35 participants were excluded. The study was approved by the Ethics Committee of the Department of Psychology, the University of the Chinese Academy of Sciences. All participants gave written informed consent.

Questionnaire assessment

The SPQ is a 74-item self-assessment scale that assesses schizotypal traits (Raine, 1991). These items can be clustered into three factors, namely the cognition-perceptual, interpersonal and disorganization factors (Chen, Hsiao, & Lin, 1997). The Chinese version of the SPQ has been shown to possess good psychometric properties (Cronbach α coefficient = 0.91, test-retest reliability = 0.82; Chen *et al.*, 1997; Wang *et al.*, 2013). The intelligence quotient (IQ) of all participants was estimated with the short-form of the Chinese version of the Wechsler Adult Intelligence Scale – Revised (Gong, 1992). The participants were asked to fill in the SPQ again before MRI scanning so that their SPQ scores were determined on the same day as the IQ estimation.

MRI acquisition

After the participants had completed the questionnaire assessment, resting-state functional magnetic resonance (rs-fMRI), anatomical and diffusion tensor images (DTI) were acquired in sequence using a 3.0 T GE MR scanner at the Institute of Psychology, the Chinese Academy of Sciences, Beijing, China from 2018 to 2019. Rs-fMRI was obtained using a T2-weighted echo planar imaging (EPI) sequence. Three hundred volumes were acquired with 37 slices, a slice thickness of 3.5 mm, a slice spacing of 0.5 mm, repetition time (TR) of 2000 ms, echo time (TE) of 30 ms, a field of view (FOV) of 220 mm, flip angle of 90°, in-plane acquisition matrix size of 64 × 64, and voxel size of 3.4 × 3.4 × 4 mm³.

The anatomical images were obtained using a T1-weighted sagittal oriented magnetization prepared rapid gradient echo (MPRAGE) sequence with the following parameters: 176 slices, a slice thickness of 1 mm, TR of 6.90 ms, TE of 2.99 ms, FOV of 256 mm, flip angle of 8°, inversion time of 450 ms, matrix size of 256 × 256, and voxel size of 1 × 1 × 1 mm³.

DTI with 64 directions were acquired with a spin echo pulse sequence: 74 slices, thickness of 2 mm, TR of 8500 ms, TE of 81 ms, FOV of 256 mm, flip angle of 90°, *b*-value of 1000 s/mm², matrix size of 128 × 128, and voxel size of 2 × 2 × 2 mm³.

DTI data preprocessing

DTI data were preprocessed by the FSL (<https://fsl.fmrib.ox.ac.uk/fsl>; Smith *et al.*, 2006) and PANDA 1.3.1 (<http://www.nitrc.org/projects/panda>; Cui, Zhong, Xu, He, & Gong, 2013). The processing was the same as our another schizophrenia DTI study (Wang *et al.*, 2020a). The data were first screened by an experienced radiologist to rule out incidental structural abnormalities. Two participants were excluded. The raw images were then cropped using the `fslroi` command of the FSL (for DWI, *f*(skull removal) of 0.25; for T1 imaging, *f*(skull removal) of 0.5; cropping gap is 3 mm). Correction of DWI eddy-current induced distortion and head motion during scanning was then carried out. DWI was registered to the *b*-value of 0 images with affine transformation (the average of these *b*-values was used as reference). Individual

images in native space were normalized to the FMRIB58_FA template (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FMRIB58_FA) in MNI space (Mori *et al.*, 2008), and smoothed using a 6 mm Gaussian kernel. Then `test_bvecs` utility was used to find the correct Invert and Swap parameter of the DWI orientation. `BedpostX` was used to build up distributions on diffusion parameters at each voxel and create all the files necessary for running the next probabilistic tractography procedure with the following parameters: fibres = 2, weight = 1, and Burn-in = 1000 (Behrens, Berg, Jbabdi, Rushworth, & Woolrich, 2007). Probabilistic tractography between 264 ROIs with 6 mm radius was performed with the following parameters: curvature threshold = 0.2, maximum number of steps = 1000, step length = 0.5 mm, number of samples = 50 000, and tracking type of path distribution (Cho *et al.*, 2015). For each ROI, probabilistic tractography was performed by seeding from all voxels in a particular ROI. The white matter connection probability from ROI *i* to ROI *j* was defined by the number of water molecules passing through ROIs divided by the total number of fibres sampled from the ROIs (Behrens *et al.*, 2007). A 264 ROIs × 264 ROIs probabilistic matrix was generated for every participant. Then a threshold value of 0.001 was applied to reduce false-positive connections between ROIs (Fang *et al.*, 2012). Since the connection probability from region *i* to region *j* was not necessarily equivalent to the one from *j* to *i*, these two probabilities were averaged to acquire a symmetrical non-directional connectivity matrix for each participant, which resulted in 34 716 connection probability values.

Rs-fMRI data preprocessing

Rs-fMRI data were preprocessed by the Statistical Parametric Mapping Software (SPM12, <https://www.fil.ion.ucl.ac.uk/spm/>) and Data Processing & Analysis for Brain Imaging (DPABI4.0, <http://www.rfmri.org/dpabi>) and brainwavelet 1.1 (<http://www.brainwavelet.org>) Software in MATLAB R2016b (MathWorks Inc., <https://www.mathworks.com>; Patel *et al.*, 2014; Yan & Zhang, 2010). The first 10 volumes were discarded. Then slice timing, realignment and wavelet despiking were performed using DPABI and brainwavelet, respectively. Subsequently, nuisance covariates, including Friston 24 head motion parameters, white matter signal and cerebrospinal fluid signal were regressed out (Friston, Williams, Howard, Frackowiak, & Turner, 1996). Structural images were manually co-registered and realigned with the anterior commissure-posterior commissure line, and co-registered to the mean functional image, and segmented as grey matter, white matter and cerebrospinal fluid using DARTEL in SPM. Then each functional image was normalized to standard Montreal Neurological Institute space in 3 × 3 × 3 mm³ voxels and the images were then band-pass filtered (0.01–0.1 Hz). Spatial smoothing was performed with a 6 mm full-width at half-maximum kernel. The mean frame-wise displacement (FD) was included as a covariate in the subsequent analysis (Power, Barnes, Snyder, Schlaggar, & Petersen, 2012; Yan *et al.*, 2013). The time series of 264 ROIs with 6 mm radius of each participant were then extracted. The resting-state fMRI networks were constructed using Pearson correlation in the DPABI 4.0, and dynamic brain networks were constructed using the dynamic conditional correlation (DCC) script in the dynamic connectivity toolbox of Martin Lindquist in MATLAB 2016b (Lindquist *et al.*, 2014; https://github.com/canlab/Lindquist_Dynamic_Correlation). A 264 ROIs × 264 ROIs static functional connectivity matrix and 264 ROIs × 264 ROIs × 290 time series DFC matrix of every

participant were generated. The mean functional connectivity strength, the DFC standard deviation and the DFC stability of every connection between two ROIs were calculated, which resulted in 34 716 values. DFC standard deviation (variability) indicates the overall fluctuation level of DFC over time. DFC stability describes the continuousness of the temporal changes in DFC between adjacent time series (Liu, Liao, Xia, & He, 2018). The formula for DFC stability is:

$$\text{DFC stability } (i, j) = 1 - \frac{1}{(T-1)} \sum_1^{T-1} \left(\frac{|r_{(i,j)(t+1)} - r_{(i,j)t}|}{2} \right)$$

where $r_{(i,j)t}$ is the strength of functional connectivity between node i and node j at time t ; T is the total number of time series: 290.

Data analysis

ANCOVA analysis of the mean functional connectivity strength, the DFC variability, the DFC stability, the white matter connectivity probability between the two groups was performed while regressing out the gender, age, IQ (FD values were also entered as a covariate in the functional connectivity analysis) based on a false discovery rate (FDR) of 0.05 using the network-based statistical software and Matlab 2016b (Verhoeven, Simonsen, & McIntyre, 2005; Zalesky, Fornito, & Bullmore, 2010). The resultant images were visualized with BrainNet Viewer (<https://www.nitrc.org/projects/bnv>; Xia, Wang, & He, 2013). Finally, Pearson correlation analysis were performed between connection probability and functional connectivity values with significant group differences and SPQ scores within the high schizotypy group while regressing out the gender, age, IQ (and FD in functional connectivity analysis) with a significance level set at FDR 0.05 correction using `fdr_bh` function in matlab 2016b (Benjamini & Hochberg, 1995).

Results

After screening and quality control, 87 individuals with high schizotypy and 122 controls were included in the final analysis. The mean age, IQ and FD of all participants was 21.44 years (s.d. = 2.18, range = 18–28 years), 124.89 (s.d. = 9.08, range = 90–145), 0.05 (s.d. = 0.02, range = 0.02–0.20) respectively. There were no significant differences between the two groups in gender, age, IQ and mean FD (38 men and 49 women, age = 21.17 ± 2.22 , IQ = 125.15 ± 9.14 , mean FD = 0.05 ± 0.03 in the high schizotypy group; 38 men and 84 women, age = 21.62 ± 2.15 , IQ = 124.72 ± 9.04 , mean FD = 0.05 ± 0.02 in the control group). The two groups differed significantly in SPQ scores (48.60 ± 5.53 in the high schizotypy group; 23.43 ± 10.62 in controls; $t = 20.22$, $p < 0.001$).

Compared with controls, the high schizotypy group exhibited increased white matter connectivity probability between the right insula (the cingulo-opercular task control network, COTCN) and the right middle frontal gyrus (the fronto-parietal task control network, FPTCN); between the left precuneus (the default mode network; DMN) and the left angular gyrus (the DMN); and increased DFC variability and decreased stability between the right precuneus (the DMN) and the left middle temporal gyrus (the DMN); as well as decreased stability between the right superior temporal gyrus (the auditory network) and the right putamen

(the subcortical network), and between the left precuneus (the DMN) and the left middle occipital gyrus (the DMN).

In addition, 18 mean functional connectivity strength values exhibited significant between-group differences. Sixteen of them were decreased in the high schizotypy group compared with controls, which mainly involved the sensorimotor network, the auditory network, the visual network, the DMN and the task control network. Increased strength values were found between the left superior frontal gyrus (the FPTCN) and the right rolandic operculum area (the auditory network), and between the right superior frontal gyrus (the DMN) and the right medial superior frontal gyrus (medsFG, the DMN; see Table 1 and Fig. 1).

Correlation analysis in the high schizotypy group found that scores on the SPQ cognition-perceptual subscale were negatively correlated with the mean functional connectivity strength between the left middle occipital gyrus (the visual network) and the left inferior parietal lobule (the FPTCN; $r = -0.32$, $p = 0.003$).

Discussion

In this study, we investigated changes in both whole-brain structural and functional connectivity in individuals with a high level of schizotypal traits. We found increased DFC variability, decreased DFC stability, and static functional connectivity strength within sub-regions of the posterior DMN and within the task control network, and decreased static functional connectivity strength between the auditory network and the task control network in the high schizotypy group. At the same time, structural connectivity between them was significantly increased in the high schizotypy group. These results support our connectivity decompensation hypothesis in individuals with high schizotypy. However, we also found increased static functional connectivity strength within the sub-regions of the anterior DMN, and between the FPTCN and the auditory network, and increased structural connectivity between the FPTCN and the auditory network in the high schizotypy group, which may indicate compensatory effects. At the same time, we found both decreased static functional connectivity and structural connectivity between the superior temporal gyrus (the auditory network) and the angular gyrus (the FPTCN). It may represent impaired neural pathways. Finally, we did not find any pathway with both increased functional connectivity and decreased structural connectivity. Our findings could extend our understanding of the temporal dynamic coupling between brain sub-regions and may provide insight into the separated large-scale brain structural and functional connectivity patterns in the high schizotypy group (Loas, Monestes, Ingelaere, Noisette, & Herbener, 2009). Moreover, our results support our hypothesis of the coexistence of 'brain connectivity decompensation' and 'brain connectivity compensation' in individuals with a high level of schizotypal traits.

Previous meta-analytic studies have reported that schizophrenia patients mainly exhibit a reduction in structural connectivity, especially at the frontal lobe and the temporal lobe (Ellison-Wright & Bullmore, 2009; Pettersson-Yeo et al., 2011). In contrast, we found that individuals with high schizotypy exhibited increased white matter connectivity probability between the right insula (the COTCN) and the right middle frontal gyrus (mFG, the FPTCN), and between the left precuneus (the DMN) and the left angular gyrus (the DMN) compared with controls. The angular gyrus and the insula integrate and connect numerous surrounding regions (Seghier, 2013; Uddin, Nomi, Hebert-Seropian, Ghaziri, & Boucher, 2017) and increased structural

Table 1. Significant white and dynamic functional connectivity differences between the high schizotypal groups and controls

ROI	SCP in high schizotypy	SCP in controls	t/p	DFC s.d. in high schizotypy	DFC s.d. in controls	t/p	DFC sta in high schizotypy	DFC sta in controls	t/p	SFC values in high schizotypy	SFC values in controls	t/p
Precentral.R(SMN,29 -17 71) -angular.R (FPTCN, 37 -65 40)	2.58E-04 (2.40E-03)	4.19E-04 (2.35E-03)	NaN	7.12E-02 (6.77E-03)	7.14E-02 (6.85E-03)	NaN	9.73E-01 (2.84E-03)	9.73E-01 (4.83E-03)	NaN	-1.43E-01 (1.47E-01)	-7.14E-02 (1.59E-01)	-12.05**
Precentral.R(SMN,20 -29 60) -iPL.L(FPTCN, -42 -55 45)	0	0	NaN	7.06E-02 (6.44E-03)	7.11E-02 (7.18E-03)	NaN	9.73E-01 (2.63E-03)	9.74E-01 (4.96E-03)	NaN	-1.17 E-01 (1.67E-01)	-3.54E-02 (1.91E-01)	-13.99**
Precentral.R(SMN,20 -29 60) -mFG.L(DMN, -35 20 51)	0	0	NaN	7.02E-02 (6.45E-03)	7.14E-02 (7.31E-03)	NaN	9.73E-01 (2.58E-03)	9.73E-01 (4.88E-03)	NaN	-1.50E-01 (1.57E-01)	-6.33E-02 (1.67E-01)	-16.95**
Precentral.L(SMN,-38 -15 69) -putamen.R (SCN, 23 10 1)	0	0	NaN	7.21E-02 (6.71E-03)	7.17E-02 (6.85E-03)	NaN	9.71E-01 (2.75E-03)	9.72E-01 (5.10E-03)	NaN	-8.78E-03 (1.22E-01)	5.35E-02 (1.28E-01)	-12.88**
SMG.R(COTCN,54 -28 34) -mFG.R(SN,26 50 27)	0	0	NaN	6.99E-02 (7.21E-03)	7.01E-02 (8.94E-03)	NaN	9.74E-01 (2.99E-03)	9.74E-01 (4.99E-03)	NaN	-7.04E-02 (2.47E-01)	6.49E-02 (2.43E-01)	-17.94**, p = 0.003
Insula.R(COTCN,36 10 1) -mFG.R(FPTCN, 32 14 56)	4.36E-03 (1.36E-02)	3.43E-04 (2.18E-03)	9.97**	7.04E-02 (6.85E-03)	7.08E-02 (7.29E-03)	NaN	9.73E-01 (2.91E-03)	9.74E-01 (4.91E-03)	NaN	-9.71E-02 (2.16E-01)	-1.00E-01 (1.99E-01)	NaN
sTG.R(AN,58 -16 7) -angular.R(FPTCN,37 -65 40)	1.19E-04 (1.11E-03)	1.12E-03 (4.32E-03)	-4.70* p = 0.03	7.10E-02 (6.72E-03)	7.20E-02 (6.36E-03)	NaN	9.73E-01 (2.72E-03)	9.73E-01 (4.91E-03)	NaN	-1.21E-01 (1.66E-01)	-2.50E-02 (1.62E-01)	-21.32**
sTG.R(AN,58 -16 7) -putamen.R(SCN,31 -14 2)	1.38E-02 (1.93E-02)	1.37E-02 (2.13E-02)	NaN	6.58E-02 (8.69E-03)	6.27E-02 (9.80E-03)	7.43* p = 0.007	9.74E-01 (3.74E-03)	9.76E-01 (5.28E-03)	-10.34**	3.56E-01 (1.97E-01)	4.30E-01 (2.20E-01)	-9.04* p = 0.003
ROA.R(AN,43 -23 20) -sFG.L(FPTCN,-23 11 64)	0	0	NaN	7.17E-02 (6.31E-03)	7.10E-02 (7.78E-03)	NaN	9.72E-01 (2.59E-03)	9.73E-01 (4.95E-03)	NaN	-2.54E-02 (1.51E-01)	-1.05E-01 (1.53E-01)	14.94**
HeschL.L(AN,-30 -27 12) -iPL.L(FPTCN,-53 -49 43)	2.48E-02 (4.08E-02)	1.46E-02 (2.44E-02)	4.87* p = 0.03	7.20E-02 (6.74E-03)	7.22E-02 (6.72E-03)	NaN	9.72E-01 (2.84E-03)	9.73E-01 (5.10E-03)	NaN	-7.80E-02 (1.37E-01)	-7.33E-03 (1.43E-01)	-13.42**
HeschL.L (AN,-30 -27 12) -iPL.L(FPTCN,-42 -55 45)	6.45E-03 (1.76E-02)	5.76E-03 (1.59E-02)	NaN	7.11E-02 (6.30E-03)	7.26E-02 (6.36E-03)	NaN	9.72E-01 (2.73E-03)	9.73E-01 (5.06E-03)	NaN	-9.85E-02 (1.53E-01)	-1.26E-02 (1.54E-01)	-17.73**
Precuneus.L(DMN,-13 -40 1) -angular.L (DMN,-39 -75 44)	4.65E-03 (1.06E-02)	1.39E-03 (5.18E-03)	9.31**	7.10E-02 (7.16E-03)	7.03E-02 (7.68E-03)	NaN	9.73E-01 (3.10E-03)	9.73E-01 (5.09E-03)	NaN	1.37E-01 (1.68E-01)	1.69E-01 (1.64E-01)	NaN
mTG.L(DMN,-46 -61 21) -precuneus.R (DMN,11 -54 17)	0	0	NaN	6.41E-02 (8.18E-03)	5.95E-02 (1.01E-02)	12.67**	9.76E-01 (3.28E-03)	9.78E-01 (5.22E-03)	-12.76**	4.22E-01 (1.77E-01)	5.39E-01 (2.17E-01)	-17.43**
sFG.R(DMN,13 55 38) -medsFG.R(DMN,6 64 22)	6.40E-02 (8.30E-02)	7.42E-02 (7.91E-02)	NaN	5.72E-02 (1.32E-02)	6.17E-02 (1.10E-02)	-8.55* p = 0.004	9.79E-01 (5.26E-03)	9.78E-01 (5.46E-03)	NaN	5.72E-01 (3.24E-01)	4.41E-01 (2.77E-01)	11.73**

aCC.R(DMN,12 36 20) -mOG.L(VN,-26 -90 3)	0	0	NaN	7.19E-02 (6.79E-03)	7.07E-02 (8.10E-03)	NaN	9.73E-01 (2.89E-03)	9.74E-01 (4.88E-03)	NaN	5.51E-02 (1.44E-01)	1.23E-01 (1.53E-01)	-12.14**
ioFG.L(DMN,-46 31 -13) -itFG.R(VAN,53 33 1)	0	0	NaN	6.36E-02 (9.67E-03)	5.89E-02 (1.16E-02)	10.56* <i>p</i> = 0.001	9.76E-01 (4.02E-03)	9.78E-01 (5.52E-03)	-12.20* <i>p</i> = 0.001	4.29E-01 (2.10E-01)	5.53E-01 (2.48E-01)	-14.83**
mOG.L(VN,-40 -88 -6) -iPL.L(FPTCN,-28 -58 48)	0	0	NaN	7.02E-02 (8.72E-03)	6.80E-02 (8.45E-03)	NaN	9.74E-01 (3.50E-03)	9.75E-01 (4.96E-03)	-6.18* <i>p</i> = 0.01	1.15E-01 (2.35E-01)	2.50E-01 (2.32E-01)	-17.77**
iOG.L(VN,-33 -79 -13) -iPL.L(FPTCN, -28 -58 48)	0	0	NaN	6.99E-02 (8.55E-03)	6.72E-02 (8.86E-03)	6.09* <i>p</i> = 0.01	9.74E-01 (3.26E-03)	9.75E-01 (5.14E-03)	-7.22* <i>p</i> = 0.008	1.58E-01 (2.48E-01)	2.99E-01 (2.39E-01)	-17.54**
Precentral.R (FPTCN,47 10 33) -iPL.R(FPTCN,49 -42 45)	2.78E-03 (1.02E-02)	1.12E-03 (5.15E-03)	NaN	6.81E-02 (1.02E-02)	6.52E-02 (1.16E-02)	NaN	9.75E-01 (3.62E-03)	9.76E-01 (5.70E-03)	-3.80* <i>p</i> = 0.05	1.86E-01 (2.98E-01)	3.30E-01 (2.91E-01)	-12.16**
Insula.L(SN,-35 20 0) -iTG.R(UN,46 -47 -17)	0	0	NaN	7.12E-02 (7.22E-03)	7.08E-02 (8.88E-03)	NaN	9.72E-01 (2.97E-03)	9.73E-01 (5.07E-03)	NaN	7.57E-02 (1.76E-01)	1.55E-01 (1.56E-01)	-13.42**
mOG.L(DMN,-41 -75 26) - precuneus.L (DMN,-11 -56 16)	1.92E-04 (1.79E-03)	9.23E-05 (1.02E-03)	NaN	6.13E-02 (9.84E-03)	5.65E-02 (1.27E-02)	8.95* <i>p</i> = 0.003	9.77E-01 (3.72E-03)	9.79E-01 (5.78E-03)	-11.62**	4.74E-01 (2.32E-01)	6.00E-01 (2.74E-01)	-11.78**

ROI, region of interest; SCP, structural connection probability; DFC, dynamic functional connectivity; SFC, static functional connectivity; s.d., standard deviation; sta, stability; NaN, not a number or un-significant; DMN, default mode network; SN, salience network; VAN, ventral attention network; FPTCN, fronto-parietal task control network; COTCN, cingulo-opercular taskcontrol network; SMN, sensorimotor network; VN, visual network; AN, auditory network; SCN, subcortical network; UN, uncertain area; medSFG, medial superior frontal gyrus; ioFG, inferior frontal gyrus, orbital part; itFG, inferior frontal gyrus, triangular part; sFG, superior frontal gyrus; mFG, middle frontal gyrus; aCC, anterior cingulate cortex; iPL, inferior parietal lobule; mOG, middle occipital gyrus; iOG, inferior occipital gyrus; sTG, superior temporal gyrus; mTG, middle temporal gyrus; iTG, inferior temporal gyrus; SMG, supramarginal gyrus; ROA, rolandic operculum area; L, left; R, right; regions are presented as name (network, peak MNI coordinates); values are presented as means (standard deviation).

**FDR 0.05 corrected; **p* < 0.05, uncorrected.

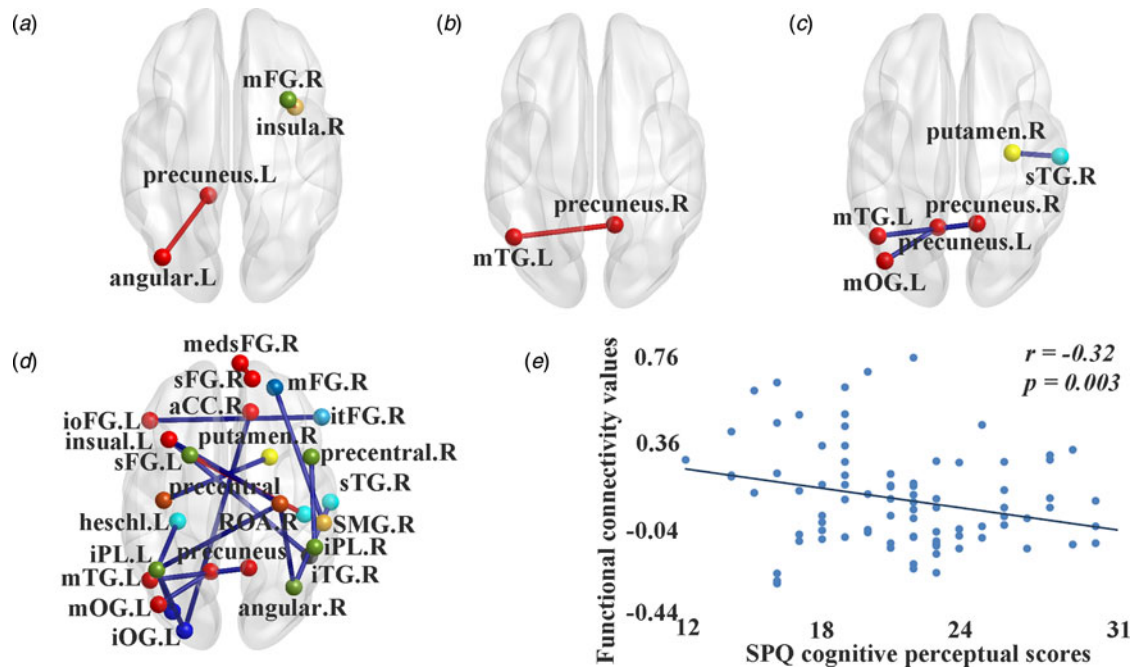


Fig. 1. Significant differences of white connectivity probability and functional connectivity values between individuals with high schizotypy and controls. (a) Significant differences in white matter connectivity probability. (b) Significant differences in dynamic functional connectivity variability. (c) Significant differences in dynamic functional connectivity stability. (d) Significant differences in static mean functional connectivity strength. (e) The negative correlation between SPQ cognitive-perceptual scores and static mean functional connectivity strength between the left middle occipital gyrus and the left inferior parietal lobule. In (a–d), red line, significant increased values in individuals with high schizotypy; blue line, significant decreased values in individuals with high schizotypy; red spot: sub-regions of the default mode network; light blue spot: sub-regions of the salience network; crimson spot: sub-regions of the sensorimotor network; blue spot: sub-regions of the visual network; green spot: sub-regions of the auditory network; cyan spot: sub-regions of the ventral attention network; lawngreen spot: sub-regions of the frontal-parietal task control network; orange spot: sub-regions of the cingulo-opercular task control network; yellow spot: sub-regions of the subcortical network; black spot: sub-regions of the uncertain area. SPQ, schizotypal personality questionnaire; medsFG, medial superior frontal gyrus; ioFG, inferior frontal gyrus, orbital part; itFG, inferior frontal gyrus, triangular part; sFG, superior frontal gyrus; mFG, middle frontal gyrus; aCC, anterior cingulate cortex; iPL, inferior parietal lobule; mOG, middle occipital gyrus; iOG, inferior occipital gyrus; sTG, superior temporal gyrus; mTG, middle temporal gyrus; iTG, inferior temporal gyrus; SMG, supramarginal gyrus; ROA, rolandic operculum area; L, left; R, right.

connectivity between these regions may represent a compensatory mechanism associated with synaptic plasticity and axonal proliferation, which mitigate dysfunctional executive control and self-sensory integration (Cai, Ryali, Chen, Li, & Menon, 2014; Nelson *et al.*, 2011; Stephan *et al.*, 2009; Zhu *et al.*, 2017). Whether these compensatory changes in white matter pathways are effective or not could be speculated from their functional connectivity changes.

However, we also found that increased DFC variability decreased DFC stability, and static functional connectivity strength within sub-regions of the posterior DMN and within the task control network, and a negative correlation between SPQ cognition-perceptual subscale scores and the mean functional connectivity strength between the sub-regions of the visual network and the task control network in the high schizotypy group. These dysfunctions may be associated with unstable brain activity, disordered self-referential thought, and impaired consciousness (Crone *et al.*, 2015; Hare *et al.*, 2018), and may be the important neuro-functional basis of abnormal perception and suspiciousness in individuals with a high level of schizotypal traits (Wong & Raine, 2018). Together with the results of increased structural connectivity between them in the high schizotypy group, this may indicate discordant brain connectivity and ineffective neurological compensation between these regions in individuals with a high level of schizotypal traits, and support the presence of connectivity decompensation within the posterior

DMN, and between the auditory network, the visual network and the task control network during the development of schizophrenia spectrum disorders.

Furthermore, the high schizotypy group also exhibited decreased DFC stability and static functional connectivity strength between sub-regions of the auditory network, the sensorimotor network and the putamen; as well as decreased static functional connectivity strength mainly between the sensorimotor network, the auditory network, the visual network, the DMN and the task control network compared with controls. The former may reflect that ineffective dopaminergic transmission for processing afferent sensory signals have already emerged in individuals with high schizotypy (Pawlak & Kerr, 2008). The latter may indicate dysfunctional segregation between auditory, visual, perceptual processing, and cognitive control ability (Carter *et al.*, 2010; Rolland *et al.*, 2015), deficits in self-awareness and external environmental information integration and perception-behaviour integration (Friston & Frith, 1995; Moore & Pope, 2014). These changes may be the key features associated with early psychosis proneness including cognitive disturbances (Kraguljac *et al.*, 2016; Wang *et al.*, 2018b; Woodward, Rogers, & Heckers, 2011), and the tendency to favour prior subjective knowledge rather than incoming sensory information (Friston *et al.*, 2016; Rolland *et al.*, 2015). Furthermore, most of these sub-regions with decreased functional connectivity features did not exhibit any changes in structural connectivity, suggesting that these

dysfunctions may be caused by grey matter neuronal dendritic changes or mediated by other connectivity (Timmler & Simons, 2019).

In addition, we found increased static functional connectivity strength within the sub-regions of the anterior DMN, mainly between the superior frontal gyrus and the medFG, and between sub-regions of the FPTCN and the auditory network, and increased structural connectivity between the FPTCN and the auditory network in the high schizotypy group. The increased connectivity between these sub-regions may indicate their self-adaptive efforts of advanced cognitive regulation for relieving sensorimotor and social-emotional dysfunction (Mohr & Claridge, 2015; Yin et al., 2011). This may be an important neurobiological compensatory mechanism for maintaining normal social behaviour in individuals with high schizotypy. The neurochemical basis of these changes may be due to the increase in functionally overlapping neural transmitters such as dopamine, glutamine, acetylcholine, and 5-hydroxytryptamine together with synaptic plasticity and axonal proliferation (Friston et al., 2016; Lau & Zuckin, 2007; Müller & Schwarz, 2007; Wolf, 2003; Wolf, Mangiavacchi, & Sun, 2003). We also found both decreased static functional connectivity and structural connectivity between the superior temporal gyrus (the auditory network) and the angular gyrus (the FPTCN) in the high schizotypy group. It may indicate defective axonal function associated with abnormal auditory sensory monitoring and self-referential processing (Cui et al., 2017; Friston et al., 2016).

Taken together, these structural and functional connectivity results may imply that enhanced connectivity within the DMN and within the task control network could not completely compensate for sensory-perceptual dysfunctional changes in individuals with a high level of schizotypal traits (Lindner, Thier, Kircher, Haarmeier, & Leube, 2005). The aberrant synaptic integration caused by hypofunction of the n-methyl-d-aspartate receptors (NMDAR) rather than axonal pathology may be the key neurochemical basis of the majority of decreased functional connectivity in individuals with high schizotypy (Tseng & O'Donnell, 2004; Wei et al., 2006), and these changes may correlate with schizophrenia genetic risk variants including DISC1, PPP1R1B, PRODH, and ZNF804A (Mothersill, Kelly, Rose, & Donohoe, 2012). Meanwhile, white matter structural alterations found in schizophrenia patients may be the pathophysiological consequence of deteriorating schizophrenia symptoms rather than the cause of the disorder (Friston et al., 2016).

This study has several limitations. First, we ascertained the absence of a psychiatric diagnosis, medication exposure, and history of psychiatric disorders of the participants only by self-report. The structured clinical interview should be adopted for higher validity in future studies. Secondly, it is difficult to clearly distinguish between compensatory and abnormal brain functional changes, and some other indirect pathways or dendritic connections between brain regions could also take effect (Timmler & Simons, 2019). Thus, our connectivity decompensation hypothesis requires further verification. Furthermore, the correlation between macroscopic connectivity and microscopic neurotransmitter and synaptic changes also requires further investigation. For example, inhibition of the dopamine D2 receptor could enhance NMDAR-dependent synaptic plasticity (Laruelle, Franke, Narendran, Kegeles, & Abi-Dargham, 2005), which may be crucial for schizophrenia pathophysiology (Abi-Saab, D'ouza, Moghaddam, & Krystal, 1998; Hahn et al., 2006). Finally, investigations on specific schizotypal dimensions or

symptoms are also important and future studies could focus on specific brain features of positive schizotypy or negative schizotypy or unusual perception or anhedonia.

In conclusion, our results suggest that individuals with high schizotypy exhibited both compensatory and deficient connectivity mainly associated with the DMN, the task control network and the sensorimotor network. Our results indicate the underlying brain connectivity adaptive changes in individuals with high schizotypy and provide a possible neurobiological basis for the connectivity decompensation hypothesis in schizophrenia spectrum disorders.

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Author contributions.

YM Wang collected, analyzed and interpreted the data, and wrote up the drafts of the manuscript. YM Wang, XL Cai, RT Zhang, YJ Zhang, HY Zhou, YY Wang, Y Wang, Y Wang, and J Huang collected the data, administered assessment and interpreted the findings. EFC Cheung interpreted the findings and commented on the drafts critically. RCK Chan generated the idea, designed the study, interpreted the findings, and commented on the drafts critically.

Conflict of interest. The authors report no conflict of interest.

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