Prefrontal-Thalamic Anatomical Connectivity and Executive Cognitive Function in Schizophrenia

Monica Giraldo-Chica, Baxter P. Rogers, Stephen M. Damon, Bennett A. Landman, and Neil D. Woodward

ABSTRACT

BACKGROUND: Executive cognitive functions, including working memory, cognitive flexibility, and inhibition, are impaired in schizophrenia. Executive functions rely on coordinated information processing between the prefrontal cortex (PFC) and thalamus, particularly the mediodorsal nucleus. This raises the possibility that anatomical connectivity between the PFC and mediodorsal thalamus may be 1) reduced in schizophrenia and 2) related to deficits in executive function. The current investigation tested these hypotheses.

METHODS: Forty-five healthy subjects and 62 patients with a schizophrenia spectrum disorder completed a battery of tests of executive function and underwent diffusion-weighted imaging. Probabilistic tractography was used to quantify anatomical connectivity between six cortical regions, including PFC, and the thalamus. Thalamocortical anatomical connectivity was compared between healthy subjects and patients with schizophrenia using region-of-interest and voxelwise approaches, and the association between PFC-thalamic anatomical connectivity and severity of executive function impairment was examined in patients.

RESULTS: Anatomical connectivity between the thalamus and PFC was reduced in schizophrenia. Voxelwise analysis localized the reduction to areas of the mediodorsal thalamus connected to lateral PFC. Reduced PFC-thalamic connectivity in schizophrenia correlated with impaired working memory but not cognitive flexibility and inhibition. In contrast to reduced PFC-thalamic connectivity, thalamic connectivity with somatosensory and occipital cortices was increased in schizophrenia.

CONCLUSIONS: The results are consistent with models implicating disrupted PFC-thalamic connectivity in the pathophysiology of schizophrenia and mechanisms of cognitive impairment. PFC-thalamic anatomical connectivity may be an important target for procognitive interventions. Further work is needed to determine the implications of increased thalamic connectivity with sensory cortex.

Keywords: Anatomical, Connectivity, Cortex, Diffusion, Schizophrenia, Thalamus

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Executive cognitive functions, which collectively encompass working memory, cognitive flexibility/set-shifting, and inhibition (1), are essential for functioning in dynamic environments. Schizophrenia is characterized by cognitive impairment, including prominent deficits in executive function (2–6). Cognitive impairment, including deficits in executive function, is an important predictor of functional outcome, making it a critical target for interventions (7). The procognitive effects of existing pharmacological and behavioral interventions are modest (8–10). The development of more effective interventions is hampered by an incomplete understanding of the neural basis of neuropsychological impairment.

Executive functions are supported by a distributed set of brain regions that includes the prefrontal cortex (PFC) and thalamus, particularly the mediodorsal (MD) nucleus, which is reciprocally connected to the PFC. Lesions to the MD nucleus often result in deficits in executive function that mimic those observed following damage to the PFC (11,12). Neuroimaging investigations indicate that the MD thalamus is a key subcortical node in a fronto-cingulo-parietal executive control network that supports working memory, cognitive flexibility, and inhibition (13). Animal models have illuminated the mechanisms of executive function. Electrophysiological investigations in rodents in particular have clarified the mechanisms of executive function by demonstrating the importance of functional coupling between the PFC and MD nucleus to working memory, attention, and cognitive flexibility (14–18).

There is compelling evidence that the PFC and thalamus are abnormal in schizophrenia. Key findings confirmed by meta-analysis include reduced gray matter volume and reduced activity during cognition (19,20). Abnormal PFC-thalamic circuitry is further supported by resting-state functional magnetic resonance imaging (fMRI) functional connectivity studies that consistently find reduced functional connectivity between the PFC and thalamus [for review, see (21)]. Several disease models are based, in part, on these findings, including the cognitive dysmetria model, which hypothesizes that the
cognitive deficits and clinical features of schizophrenia result from a core defect in cortical-subcortical circuitry (22–26).

Regardless of the model, critical knowledge gaps remain. First, while altered PFC-thalamic activity and functional connectivity is well established, considerably less is known about the integrity of anatomical connectivity between the PFC and thalamus. A small number of diffusion-weighted imaging studies have examined thalamocortical anatomical connectivity; however, they have yielded inconclusive results, likely owing to small sample sizes in some cases (27–29). Second, the cognitive correlates of PFC-thalamic structural connectivity, especially executive function, are poorly understood. Finally, the integrity of structural connections linking other cortical areas to the thalamus has received little attention. The current study was undertaken to address these knowledge gaps and limitations of prior studies. The primary aim of this investigation was to test the hypothesis that PFC–MD thalamic anatomical connectivity is 1) reduced in schizophrenia and 2) related to deficits in executive function. A secondary aim of this investigation was to characterize connectivity disturbances in other anatomical networks linking the cortex to thalamus.

**METHODS AND MATERIALS**

**Study Participants**

Forty-seven healthy subjects and 67 patients with a schizophrenia spectrum disorder (i.e., schizophreniform disorder, schizophrenia, schizoaffective disorder) who participated in an ongoing study of thalamocortical networks in psychotic disorders were screened for inclusion in this study. All patients with a schizophrenia spectrum diagnosis, hereafter referred to as schizophrenia, were recruited through the Psychotic Disorders Program at the Vanderbilt Psychiatric Hospital. Healthy subjects were recruited from Nashville and the surrounding area via advertisement and word of mouth. This study was approved by the Vanderbilt University Institutional Review Board, and all subjects provided written informed consent before participating.

Psychiatric diagnoses were confirmed in patients and ruled out in healthy subjects using the Structured Clinical Interview for DSM (30). The Positive and Negative Syndrome Scale (PANSS) (31) was administered to patients to quantify the severity of clinical symptoms of psychosis. The Wechsler Test of Adult Reading (32) was administered to all subjects to provide an estimate of premorbid IQ. Exclusion criteria included age younger than 16 years or older than 55 years; estimated premorbid IQ less than 70; presence of a systemic medical illness (e.g., diabetes, cardiovascular disease) or central nervous system disorder (e.g., multiple sclerosis, epilepsy) that would affect study results; history of significant head trauma; psychotropic drug use (healthy subjects only); active substance abuse, based on Structured Clinical Interview for DSM criteria, within the past 3 months (or lifetime history of substance abuse and/or dependence in healthy subjects); and MRI contraindications (e.g., metal implants, claustrophobia).

**Neuropsychological Testing**

All study participants completed a brief neuropsychological assessment. Given our a priori focus on executive function and PFC-thalamic circuitry, testing focused on assessing the three domains of executive function identified by Miyake et al. (1): working memory, set-shifting/cognitive flexibility, and inhibition. The Wechsler Memory Scale–Third Edition (WMS-III) (33) digit and spatial span subtests, which comprise the Working Memory Index; Wisconsin Card Sorting Test (WCST) 64 Card Version (34); and Continuous Performance Test–AX version (AX-CPT) (35) were administered to quantify working memory, set-shifting, and inhibition, respectively. The WMS-III Working Memory Index, WCST total errors, and d-prime measure from the AX-CPT served as the dependent variables. The Screen for Cognitive Impairment in Psychiatry (36) was also administered to quantify overall neuropsychological functioning.

**Neuroimaging Data Acquisition, Preprocessing, and Probabilistic Tractography**

Neuroimaging data were acquired on a 3T Philips Intera Achieva scanner (32-channel receive head coil, single-band imaging; Philips Healthcare, Andover, MA) located at the Vanderbilt University Institute for Imaging Sciences. A T1-weighted structural scan (1-mm isotropic resolution) and high-angular radial diffusion-weighted imaging (HARDI) scan (2.5-mm isotropic resolution, 60 directions, b value = 2000 s/mm, 5 b0 images) were collected for each subject in a single imaging session. Of note, the HARDI data were collected with a sensitivity encoding factor of 2.2 to reduce echo time and echo-planar image distortions.

Each subject’s structural T1-weighted MRI was automatically segmented using the program Multi Atlas developed by one of the authors (BAL) (37). Briefly, Multi Atlas uses a statistical fusion framework built on an a priori model comprising a manually traced dataset to automatically label 133 cortical and subcortical brain structures. Selected cortical structures were combined to generate six bilateral cortical regions of interest (ROIs) that along with the thalamus were used as targets and the seed, respectively, for probabilistic tractography. The six cortical ROIs corresponded to the PFC, motor cortex/supplementary motor area, somatosensory cortex, posterior parietal cortex, temporal cortex, and occipital cortex (see the Supplement for an example segmentation and list of cortical structures included in each cortical ROI). The thalamus ROI was manually edited by one author (author MG-C), blinded to diagnostic status, to include the lateral and medial geniculate nuclei. In addition, each subject’s T1-weighted anatomical image was segmented into gray matter, white matter, and cerebrospinal fluid using the VBM8 toolbox (http://www.neuro.uni-jena.de/vbm/download/), which employs the DARTEL algorithm for high-dimensional spatial normalization to Montreal Neurological Institute (MNI) space (38). Before analysis, all HARDI data underwent quality assurance (QA) using an automated QA pipeline developed by our group that includes visualization of HARDI data and extraction of mean head translation and rotation in the X, Y, and Z directions and average number of voxels (in percent) rejected per gradient owing to poor diffusion tensor imaging model fitting (39).

Following QA, HARDI data preprocessing and probabilistic tractography were performed using FMRIB’s Diffusion Toolbox for FSL version 5.0.6 software package (http://www.fmrribox.ac.uk/fsl/). Data preprocessing and probabilistic tractography were performed using FMRIB’s Diffusion Toolbox for FSL version 5.0.6 software package (http://www.fmrribox.ac.uk/fsl/). Data preprocessing and probabilistic tractography...
are described in detail in the Supplement. Briefly, seed-to-target tractography analyses were run using probtrackx2 [40] to quantify anatomical connectivity between the thalamus (i.e., seed) and each of the six cortical ROIs (i.e., targets). The left and right hemispheres were analyzed separately. From each thalamic voxel, 5000 samples were sent through the probability distributions on principal fiber direction. From probtrackx2, seed-to-target voxelwise images were generated in which the value of each voxel within the seed mask (i.e., thalamus) represents the number of samples seeded from that voxel reaching the relevant target mask. The connectivity of each cortical region with the thalamus was calculated by dividing the number of samples reaching that region, summed across all voxels in the thalamus, by the total number of samples within the thalamus reaching all cortical regions [29]. This is a measure of total tractography-defined connectivity from the thalamus to a particular cortical area, independent of where the tract originated from inside the thalamus and after controlling for overall connectivity of the thalamus. These values, expressed as total connectivity (in percent), served as the dependent variables for the primary analysis described below. In addition, voxelwise probability maps were generated by dividing the number of samples from each voxel that arrived at the corresponding target by the total number of samples from that voxel reaching any cortical region. These probability maps were used in the voxelwise analysis described below.

**Statistical Analysis**

Group differences in dichotomous and continuous demographic and cognitive variables were examined using χ² and independent groups t tests, respectively. For the analysis of total cortical-thalamic connectivity, left and right hemisphere connectivity values were averaged, as we did not have any a priori hypotheses regarding laterality and to maximize statistical power. This resulted in six dependent variables per subject, one for each cortical target ROI. Total percent connectivity values were analyzed using independent groups t tests. Given our a priori hypothesis that PFC-thalamic connectivity would be reduced in schizophrenia, no correction was applied to the critical α for this contrast. Significance was set to p = .01 (i.e., Bonferroni corrected) for the remaining five cortical target ROIs. The primary analysis described above was complemented with a voxelwise analysis to further localize potential group differences in thalamocortical structural connectivity (see Supplement). Voxelwise analyses were thresholded at the cluster-level Pfamilywise error [FWE] corrected = .05 for voxelwise Puncorrected = .001. Finally, to test the hypothesis that PFC-thalamic anatomical connectivity is related to cognitive impairment in schizophrenia, average PFC-thalamic connectivity was correlated with each measure of executive function (WMS-III Working Memory Index, WCST total errors, AX-CPT d-prime) using partial correlation analysis controlling for age and sex. Similarly, average connectivity was extracted from significant clusters identified in the PFC-thalamic voxelwise analyses and correlated with each of the executive cognitive measures using partial correlation analysis controlling for age and sex.

**RESULTS**

Seven subjects (2 healthy individuals and 5 patients) were excluded owing to presence of neuroimaging data artifacts based on visual inspection. Thus, the final sample included 45 healthy subjects and 62 patients with schizophrenia (Table 1). With the exception of education, the groups did not differ in demographic variables. No significant differences in HARDI QA metrics were detected between groups, including mean head

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**Table 1. Sample Demographics**

<table>
<thead>
<tr>
<th></th>
<th>Healthy Subjects (n = 45)</th>
<th>Schizophrenia (n = 62)</th>
<th>Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex, Male/Female</td>
<td>29/16</td>
<td>43/19</td>
<td>0.29</td>
</tr>
<tr>
<td>Race, White/Black/Other</td>
<td>32/9/4</td>
<td>44/16/2</td>
<td>1.87</td>
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<tr>
<td>Age, Years</td>
<td>27.8 (7.0)</td>
<td>27.0 (6.6)</td>
<td>0.53</td>
</tr>
<tr>
<td>Education, Years</td>
<td>16.3 (2.0)</td>
<td>13.6 (2.1)</td>
<td>6.96</td>
</tr>
<tr>
<td>Parental Education, Years</td>
<td>15.0 (2.1)</td>
<td>14.8 (2.9)</td>
<td>0.45</td>
</tr>
<tr>
<td>Premorbid IQ</td>
<td>113.0 (10.2)</td>
<td>102.6 (13.8)</td>
<td>4.24</td>
</tr>
<tr>
<td>SCIP Z Score</td>
<td>0.39 (0.56)</td>
<td>-0.73 (0.77)</td>
<td>8.30</td>
</tr>
<tr>
<td>WMS-III WM Index</td>
<td>107.5 (12.7)</td>
<td>95.0 (12.1)</td>
<td>5.16</td>
</tr>
<tr>
<td>WCST Total Errors</td>
<td>11.1 (6.6)</td>
<td>19.6 (10.6)</td>
<td>4.62</td>
</tr>
<tr>
<td>AX-CPT D-Prime</td>
<td>3.54 (0.63)</td>
<td>2.70 (1.15)</td>
<td>4.47</td>
</tr>
<tr>
<td>Age at Illness Onset, Years</td>
<td>21.8 (5.7)</td>
<td>5.7 (6.4)</td>
<td></td>
</tr>
<tr>
<td>Duration of Illness, Years</td>
<td>5.7 (6.4)</td>
<td>5.7 (6.4)</td>
<td></td>
</tr>
<tr>
<td>APD Dose, CPZ Equivalents, mg/day</td>
<td>353.6 (224.0)</td>
<td>13.7 (7.2)</td>
<td></td>
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<tr>
<td>PANSS Positive</td>
<td>14.2 (5.7)</td>
<td>14.2 (5.7)</td>
<td></td>
</tr>
<tr>
<td>PANSS General</td>
<td>27.4 (7.4)</td>
<td>27.4 (7.4)</td>
<td></td>
</tr>
</tbody>
</table>

Values are presented as n or mean (SD).

between-group analyses revealed several differences in connectivity between healthy subjects and patients with schizophrenia (Figure 2D). Consistent with our hypotheses and the primary analysis presented above, anatomical connectivity between the thalamus and PFC was decreased in schizophrenia in two clusters located in the MD thalamus at MNI coordinates $-6, -18, 10$ (cluster size = 24 voxels, $p_{FWE-corrected} = .022$) and $-6, -12, 2$ (cluster size = 18 voxels, $p_{FWE-corrected} = .028$). To determine what areas of the PFC these regions are connected to, we performed a separate probabilistic tractography analysis using these clusters as seeds and the PFC as the target ROI (see Supplement). This analysis revealed that the two thalamic clusters are connected to the lateral PFC, including superior, middle, and inferior frontal gyri (see Supplement). Qualitatively, there was less overlap across patients in the schizophrenia group in the paths linking PFC to the two thalamic clusters.

Additionally, the voxelwise analysis revealed group differences in thalamic connectivity with motor cortex, sensorimotor cortex, posterior parietal cortex, and occipital lobe. Specifically, clusters of both decreased and increased connectivity in schizophrenia were detected for motor-thalamic connectivity, whereas increased thalamic connectivity was observed for somatosensory, posterior parietal, and occipital cortex (Figure 2D). Voxelwise results are detailed in the Supplement.

**PFC-Thalamic Anatomical Connectivity and Cognitive Impairment in Schizophrenia**

The patients with schizophrenia performed worse than healthy subjects on all tests of executive function (Table 1). To test the hypothesis that impaired executive function in schizophrenia is related to PFC-thalamic anatomical connectivity, the three executive function measures (WMS-III Working Memory Index, WCST total errors, and AX-CPT d-prime) were correlated with total PFC-thalamic connectivity, and average PFC-thalamic connectivity was extracted from the two clusters identified in the voxelwise analysis. None of the measures of executive cognitive function correlated with total PFC-thalamic connectivity.

**Thalamocortical Structural Networks in Schizophrenia**

Voxelwise analyses are presented in Figure 2. As shown in Figure 2B and C, the connectivity patterns appeared qualitatively similar in both groups and consistent with the anatomy of the thalamus and prior neuroimaging investigations (41,42). Each cortical region was anatomically connected to distinct, largely nonoverlapping regions of the thalamus. The translation and rotation in X, Y, and Z directions and percentage of outlier voxels per gradient with poor fitting diffusion tensor imaging (see Supplement). All but 12 patients were taking an antipsychotic medication. Antipsychotic dosage was unavailable for 2 patients.

**Thalamocortical Anatomical Connectivity**

Total percent connectivity of each cortical ROI with the thalamus in healthy individuals and patients with schizophrenia is presented in Figure 1. Complete statistical results are presented in the Supplement. Consistent with our hypothesis, anatomical connectivity between the PFC and thalamus was reduced in schizophrenia ($t_{105} = 3.06, p = .003$). In contrast to the reduction in PFC-thalamic connectivity, patients with schizophrenia demonstrated significantly increased thalamic connectivity with somatosensory ($t_{105} = 2.69, p = .008$) and occipital cortices ($t_{105} = 2.80, p = .006$). Follow-up analyses revealed that PFC-thalamic anatomical connectivity was lower in both left and right hemispheres, although only the left hemisphere reached statistical significance ($t_{105} = 3.73, p < .001$). Increased thalamic connectivity with the occipital cortex was also more prominent in the left hemisphere, whereas somatosensory connectivity was increased bilaterally. Of note, total volume of the thalamus and cortical ROIs used as seed and target masks for probabilistic tractography did not differ between groups (see Supplement). Additionally, average daily dose of antipsychotic medication (in chlorpromazine equivalents) was unrelated to thalamocortical connectivity (all $r_s < .22, p > .142$).

**Thalamocortical Anatomical Connectivity: Voxelwise Analysis**

Voxelwise analyses are presented in Figure 2. As shown in Figure 2B and C, the connectivity patterns appeared qualitatively similar in both groups and consistent with the anatomy of the thalamus and prior neuroimaging investigations (41,42). Each cortical region was anatomically connected to distinct, largely nonoverlapping regions of the thalamus. The
connectivity. Similarly, none of the measures of executive function correlated with PFC-thalamic total connectivity in healthy subjects (1 healthy subject who had an exceptionally high WMS-III Working Memory Index score of 151 was excluded from the analysis).

With respect to the voxelwise analysis, neither of the two thalamic clusters that demonstrated reduced connectivity with the PFC was related to impaired AX-CPT and WCST performance in patients. However, average connectivity within one of the clusters (MNI = −6, −18, 10) positively correlated with WMS-III Working Memory Index (r = .30, p = .020), indicating that higher PFC-thalamic structural connectivity was associated with better working memory (Figure 3). The same correlation did not reach significance in healthy subjects (r = −.05, p = .778). A regression analysis with age, sex, group, and group × PFC connectivity (at MNI = −6, −18, 10) interaction term entered as predictors of WMS-III Working Memory Index revealed that the association between working memory and connectivity at the MNI cluster = −6, −18, 10 did not differ between groups (group × connectivity interaction term: t(104) = 1.41, p = .161). PFC connectivity at the other cluster identified in the between-groups analysis (MNI = −6, −12, 2) was unrelated to WMS-III Working Memory Index in both patients with schizophrenia and healthy subjects (partial r < .18, p > .261).

To assess the relative specificity of the association between PFC-thalamic connectivity and cognition in schizophrenia for the cluster located at MNI = −6, −18, 10, we examined the correlation between Screen for Cognitive Impairment in Psychiatry scores and average connectivity within this cluster. The Screen for Cognitive Impairment in Psychiatry is composed of several subtests, including a version of the Auditory Consonant Trigrams test of working memory, a word list learning test of verbal learning and delayed recall, phonemic verbal fluency, and a measure of psychomotor processing speed [see (36,43)]. Of these subtests, only the verbal working memory subset correlated with PFC-thalamic anatomical connectivity in the schizophrenia group (partial r = .27, p = .036; all remaining partial r’s < .14, p > .304).

Finally, to explore the potential cognitive correlates of hyperconnectivity, average connectivity was extracted from the clusters that demonstrated hyperconnectivity with motor, somatosensory, posterior parietal, and occipital cortex in the between-groups voxelwise analysis and correlated with executive function measures. This analysis revealed that occipital hyperconnectivity with the cluster located at MNI = −8, −28, 2 inversely correlated with AX-CPT d-prime (r = −.31, p = .018), and somatosensory hyperconnectivity with the thalamic cluster located at MNI 12, −20, 6 correlated positively with WMS-III Working Memory Index (r = .27, p = .042).

**Thalamocortical Anatomical Connectivity and Clinical Symptoms in Schizophrenia**

Although no relationships were hypothesized a priori, associations between thalamocortical anatomical connectivity and
clinical symptoms in patients (i.e., PANSS positive, negative, and general scores) were examined using partial correlations controlling for age and sex. Total connectivity measures were unrelated to PANSS scores (all partial rs < .24, p > .060). With respect to the voxelwise analysis, both thalamic clusters that demonstrated reduced connectivity with the PFC in schizophrenia correlated inversely with PANSS general symptoms indicating that severity of general symptoms was related to lower PFC-thalamic connectivity (cluster at MNI −6, −12, 2: r = −.32, p = .012; cluster at MNI −6, −18, 10: r = −.27, p = .039). Additionally, hyperconnectivity of the occipital cortex at the thalamic cluster located at MNI −8, −28, 8 and motor cortex at the thalamic cluster located at MNI −8, −6, 0 positively correlated with general symptoms (r = .29, p = .027) and positive symptoms (r = .26, p = .043), respectively.

**DISCUSSION**

Motivated by evidence implicating PFC-thalamic circuitry in the pathophysiology of schizophrenia and mechanisms of cognitive impairment, we tested the hypothesis that anatomical connectivity between the PFC and thalamus is 1) reduced in schizophrenia and 2) related to impaired executive function. Our first hypothesis was supported; total connectivity of the PFC with the thalamus was significantly reduced in schizophrenia. Voxelwise analyses confirmed the reduction in connectivity localized to the MD aspect of the thalamus where paths linking the thalamus to lateral PFC originate from. Our second hypothesis was partially supported. While total PFC connectivity with the thalamus did not correlate with any measure of executive function, average connectivity within a cluster located in the MD thalamus that demonstrated reduced connectivity in patients correlated with working memory. Similar associations were not found for other measures of executive function and overall neuropsychological functioning.

Our findings add to the modest literature on thalamocortical anatomical connectivity in schizophrenia that so far has produced mixed findings. Our results are consistent with a prior study of 15 patients with schizophrenia by Marenco et al. (29), which, using almost identical methods, found that PFC-thalamic connectivity is reduced in schizophrenia. Marenco et al. (29) also reported a correlation between lateral PFC-thalamic connectivity and working memory in a subset of nine patients that, while not statistically significant, was very similar to the current findings (r = .38 vs. r = .30). However, both studies are at odds with two recent investigations that found reduced connectivity with the orbitofrontal cortex but not the lateral and medial PFC (27,28). The inconsistent findings may be due to differences in sample sizes, as the current investigation included substantially more patients than prior studies; differences across studies in the methods used to quantify anatomical connectivity and define cortical targets; difficulties imaging the orbitofrontal cortex and possibly reduced reliability of orbitofrontal cortex–thalamic connectivity [e.g., see (29)]; and heterogeneity across patient samples. Interestingly, average connectivity within the two clusters in the thalamus that demonstrated reduced PFC-thalamic connectivity in schizophrenia also correlated with severity of general clinical symptoms. This finding was not hypothesized a priori and should be interpreted cautiously given the exploratory nature of the analysis. Nonetheless, it suggests that PFC-thalamic pathology may underlie some dimensions of clinical symptoms and may also account for the heterogeneous findings across studies.

The results support thalamocortical network models of schizophrenia and cognitive impairment (22,24,26,44,45). They are also consistent with functional imaging studies, particularly
resting-state fMRI investigations, that consistently find reduced functional connectivity between the PFC and thalamus in schizophrenia [reviewed in (21)]. Whether disrupted connectivity results from pathology in the PFC and/or thalamus is an unresolved question that is especially difficult to address with neuroimaging owing to the reciprocal nature of anatomical connections between the thalamus and cortex. Recent electrophysiology studies in rodents highlight the importance of resolving this question, not only for understanding the nature of cognitive impairment in schizophrenia but also for developing procognitive interventions. Specifically, Bolkan et al. (15) found that MD thalamus–PFC coupling is essential for sustaining working memory delay-related activity in the PFC, whereas PFC-thalamic functional coupling is critical during the choice selection phase of working memory. Similarly, Schmitt et al. (18) found that MD thalamus stimulation enhances functional connectivity within the PFC and improves performance on forced-choice tests of attentional control, in contrast to direct stimulation of the PFC, which impairs task performance. Combined, the results indicate that pathology within specific nodes of the executive control network may lead to differential cognitive impairment and, by extension, suggest that targeting different nodes may improve specific aspects of behavior. Interestingly, a recent clinical trial found that cognitive remediation training improves PFC-thalamic functional connectivity, which in turn correlates with the degree of improvement in overall cognitive functioning (46); these findings are consistent with the broader literature on the effect of cognitive remediation training on PFC and thalamic activity (47). While agnostic with respect to the primary site of dysfunction (i.e., PFC, thalamus, or both), these findings provide compelling evidence that PFC-thalamic circuitry may be an important target for improving cognitive function in schizophrenia.

While the focus of our study was on PFC-thalamic connectivity, we also examined the integrity of thalamic connections with other cortical areas. In contrast to the PFC, anatomical connectivity with other cortical areas, primarily somatosensory and occipital cortex and to a lesser extent motor and posterior parietal cortex, was increased in schizophrenia. Again, the findings are reminiscent of resting-state fMRI investigations, which consistently find that PFC-thalamic hypoconnectivity in schizophrenia is accompanied by thalamic hyperconnectivity with sensory and motor cortical areas (48–59). However, in contrast to functional disconnection, which could result from alterations in direct or indirect connections, abnormalities in connectivity inferred from diffusion-weighted imaging presumably reflect disruption of direct anatomical connections (60). It is possible that anatomical hyperconnectivity reflects novel anatomical connections and/or a failure to prune some connections during development. In humans, segregation of neurons into thalamic nuclei begins at 10 to 14 weeks of gestation and continues until 40 weeks (61,62). The development of thalamocortical anatomical connections is similarly prolonged and relies on a complex interplay between genes and intrinsic neuronal signaling (63,64). Consequently, alterations in the expression and timing of certain genes, possibly in combination with extrinsic factors, could theoretically lead to abnormalities in the patterning of connections between the cortex and the thalamus. Very little is known about the postnatal development of thalamic nuclei and anatomical connectivity. The limited available evidence suggests that connections between the thalamus and association cortical areas, especially the PFC, mature later than connections linking the thalamus with primary sensory and motor areas (65,66). Interestingly, some of the thalamic regions that demonstrated hyperconnectivity with areas outside the PFC correlated with severity of clinical symptoms and cognitive impairment. While these findings suggest there may be functional consequences of anatomical hyperconnectivity and therefore may merit further investigation, they should also be interpreted cautiously given the lack of a priori hypotheses and the number of correlations performed.

It is also possible that the abnormalities in anatomical connectivity reflect differences in the relative volume of thalamic nuclei and/or disruption of postnatal brain development. For instance, the combination of reduced PFC-thalamic and elevated sensory-thalamic anatomical connectivity may result from relative volume loss of the MD nucleus and expansion of ventral posterior nuclei and pulvinar. Reduced thalamic volume is a consistent finding in clinical neuroimaging studies (19); however, most conventional structural imaging methods are not able to delineate thalamic nuclei owing to lack of contrast and, in some cases, limited resolution (67). One exception is proton density imaging, which can be used to resolve the lateral and medial geniculate nuclei. Unfortunately, we did not collect proton density images, which is an important limitation of the study given our finding of increased occipital-thalamic connectivity in schizophrenia. Similarly, it is also important to note that diffusion-weighted imaging is an indirect method for inferring white matter integrity and that the reliability of thalamocortical connectivity derived from probabilistic tractography was not confirmed in the current investigation. However, prior studies suggest that test-retest reliability of thalamocortical connectivity is reasonable and probabilistic tractography is useful for tracking disease progression in white matter (29,68).

In conclusion, PFC-anatomical connectivity is reduced in schizophrenia and correlated with the severity of working memory impairment. In addition to illuminating the pathophysiology of schizophrenia, the present results suggest that PFC-thalamic connectivity may be a useful neural target for procognitive interventions.

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Thalamocortical Structural Networks in Schizophrenia

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