Impaired cerebellar functional connectivity in schizophrenia patients and their healthy siblings

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The long-standing notion of schizophrenia as a disorder of connectivity is supported by emerging evidence from recent neuroimaging studies, suggesting impairments of both structural and functional connectivity in schizophrenia. However, investigations are generally restricted to supratentorial brain regions, thereby excluding the cerebellum. As increasing evidence suggests that the cerebellum contributes to cognitive and affective processing, aberrant connectivity in schizophrenia may include cerebellar dysconnectivity. Moreover, as schizophrenia is highly heritable, unaffected family members of schizophrenia patients may exhibit similar connectivity profiles. The present study applies resting-state functional magnetic resonance imaging to determine cerebellar functional connectivity profiles, and the familial component of cerebellar connectivity profiles, in 62 schizophrenia patients and 67 siblings of schizophrenia patients. Compared to healthy control subjects, schizophrenia patients showed impaired functional connectivity between the cerebellum and several left-sided cerebral regions, including the hippocampus, thalamus, middle cingulate gyrus, triangular part of the inferior frontal gyrus, supplementary motor area, and lingual gyrus (all p < 0.0025, whole-brain significant). Importantly, siblings of schizophrenia patients showed several similarities to patients in cerebellar functional connectivity, suggesting that cerebellar dysconnectivity in schizophrenia might be related to familial factors. In conclusion, our findings suggest that dysconnectivity in schizophrenia involves the cerebellum and that this defect may be related to the risk to develop the illness.

Keywords: cerebellum, schizophrenia, siblings, functional connectivity, resting-state fMRI, dysconnectivity

INTRODUCTION
The cerebellum has long been regarded as a brain structure that is exclusively involved in motor systems, but an increasing body of evidence suggests that it is also involved in cognition and emotion (Schmahmann and Caplan, 2006). The cerebellum is thought to influence motor systems by evaluating disparities between intention and action and by adjusting the operation of motor cortices accordingly, through feed-back and -forward loops via the thalamus and pons (Kandel et al., 2000). The cerebellum may be integrated in the neural circuits governing higher cognitive functions in a similar fashion. Cerebellar modulation of cognitive processes was shown in error-related learning and timing, but a more general involvement has also been suggested (Schmahmann, 2006; Andreasen and Pierson, 2008). According to the “cognitive dysmetria” and “dysmetria of thought” models of schizophrenia – a severe psychiatric disorder characterized by hallucinations, delusions, and disintegration of thinking – aberrant cerebellar modulation of information from and to the cerebral cortex may be a part of the pathophysiology of schizophrenia (Andreasen et al., 1998; Schmahmann, 1998).

The notion that schizophrenia involves the aberrant integration of information between anatomically separated brain regions is long-standing (Wernicke, 1906; Bleuler, 1911; Kraepelin, 1919; Friston, 1998; Stephan et al., 2009) and is supported by recent neuroimaging studies showing aberrant structural (Assaf and Pasternak, 2008; Bassett et al., 2008; Bassett and Bullmore, 2009; Van den Heuvel et al., 2010; Zalesky et al., 2011) and functional (Lynall et al., 2010; Fornito et al., 2011) connectivity in schizophrenia (Pettersson-Yeo et al., 2011; Rubinov and Bassett, 2011). However, the cerebellum is typically excluded from these “whole-brain” analyses of brain connectivity in schizophrenia. Interestingly, some studies using diffusion tensor imaging (DTI) to target specific white matter tracts have shown impaired structural connectivity of the cerebellum (Kanaan et al., 2009; Kyriakopoulos and Frangou, 2009). Furthermore, a few functional connectivity studies in schizophrenia have reported impaired functional integration of the cerebellum (Honey et al., 2005; Kim et al., 2008; Becerril et al., 2011; Repovs et al., 2011). However, it remains unclear whether functional connectivity between the cerebellum and the rest of the brain is affected in schizophrenia, and if so, to what extent. Functional connectivity between anatomically separated brain regions is defined as the temporal dependency of their neural activation patterns and is thought to be reflected, to some extent, by their coherence in spontaneous (resting-state) fluctuations in functional magnetic resonance imaging (fMRI) signal (Friston et al., 1993; Biswal et al., 1995, 1997; Van den Heuvel and Hulshoff Pol, 2010).
A total of 62 schizophrenia patients, 67 siblings of schizophrenia patients, and 41 healthy comparison subjects were included in this study. Age at time of scanning, presence or absence of current and lifetime psychopathology was established for all participants, using the Comprehensive Assessment of Symptoms and History (CASH) interview (Andreasen et al., 1992). Schizophrenia patients were eligible for the present study if they met Diagnostic and Statistical Manual of Mental Disorders fourth edition (DSM IV; American Psychiatric Association, 1994) criteria of schizophrenia or related spectrum disorders. Both siblings of schizophrenia patients and healthy comparison subjects could have no history of any psychiatric illness, including substance dependence of abuse; and healthy comparison subjects had no first- or second-degree family members with a lifetime psychotic disorder.

Study participants originated from a total number of 132 families. Within the healthy comparison group, there were two sibling pairs (both healthy control subjects, i.e., without a first- or second-degree relative with a lifetime psychotic disorder), and within the 67 siblings of patients, there were seven pairs of siblings of a schizophrenia patient and one set of three siblings of a schizophrenia patient. There were no family relationships within the patient group. Between the sibling and patient groups, there were a total number of 27 family-ties [i.e., a schizophrenia patient and (one of) their sibling(s) participated in the study].

Presence or absence of current and lifetime psychopathology was assessed using the positive and negative syndrome scale (PANSS; Kay et al., 1987); Furthermore, the type and daily dose of antipsychotic medication at the time of scanning was recorded, and a haloperidol equivalent dose was calculated using conversion rates (risperidone 0.5:1; olanzapine 1.66:1; quetiapine 25:1; clozapine 33.33:1; aripiprazole 2.5:1; flupenthixol 0.66:1; perphenazine 2.66:1; Kroken et al., 2009). In healthy comparison subjects and siblings of patients, a shortened version of the structured interview of schizotypy-revised (SIS-R; Kendler et al., 1989; Vollema and Ormel, 2000) was employed [excluding four items: social isolation (last 3 years); antisocial behavior; dysfunction (obligatory activities); and dysfunction (personal caretaking)], to measure the overall severity of schizotypal signs and symptoms on a four-point scale (0 = absent, 1 = mild, 2 = moderate, 3 = severe). For all participants, global cognitive functioning, as measured by total IQ, was estimated using four subtests (i.e., Information, Arithmetic, Block design, and Digit symbol coding) of the Dutch version of the Wechsler adult intelligence scale (WAIS; Stinissen et al., 1970).

Finally, statistical testing of group-differences in demographic characteristics was performed using analysis of variance (ANOVA) for continuous and Chi-Square tests for categorical variables. All demographic and clinical characteristics are provided in Table 1.

### MATERIALS AND METHODS

#### PARTICIPANTS

A total of 62 schizophrenia patients, 67 siblings of schizophrenia patients, and 41 healthy comparison subjects were included in this study. Study participants were recruited at the University Medical Center Utrecht, during a large ongoing cohort in the Netherlands (Genetic Risk and Outcome of Psychosis; GROUP). The study was approved by the affiliated ethical committee. Study participants were between 18 and 60 years of age. All subjects provided written informed consent prior to participation. Subjects with a history of head trauma or major medical or neurological illness were excluded.

#### IMAGE ACQUISITION AND PREPROCESSING

Resting-state fMRI data were acquired on two 1.5 T Magnetic Resonance Imaging scanners (Philips Medical Systems, Best, The Netherlands) at the University Medical Center Utrecht, The Netherlands. BOLD time-series were recorded during 9 minutes using a 3D-PRESTO sequence (Van Gelderen et al., 1995; Ramsey et al., 1996; acquisition parameters: TR/TE 21.1/31.1 ms; voxel size 4 mm × 4 mm × 4 mm). Subsequently, a T1 weighted image was acquired for anatomical reference (3D FFE pulse sequence, TR/TE = 30/4.6 ms, flip-angle 30°, FOV 256 mm × 256 mm, voxel size 1 mm × 1 mm × 1.2 mm, 160–180 contiguous slices; Ramsey et al., 2006). Preprocessing was performed using SPM5 software. In short, all resting-state functional images were registered to the last functional scan to correct for head movements and co-registered with the T1 scan to ensure overlap between the anatomical reference scan and resting-state time-series. The registered functional images were spatially smoothed, using an 8-mm full width half-max smoothing kernel. Next, the T1 scan and resting-state time-series were normalized to standard space, matching the MNI-152 template. To create individual anatomical label maps, the spatially normalized T1 image was overlaid with the automated anatomical labeling (AAL) template (Tzourio-Mazoyer et al., 2002), distinguishing 116 anatomical brain regions. Time-series were corrected for white matter, ventricle, and motion parameters (using regression) and band pass filtered (bandwidth 0.01–0.1 Hz) to eliminate low frequency noise and influences of frequencies reflecting possible cardiac or respiratory oscillations.

### CEREBELLAR REGION OF INTEREST DEFINITION

The cerebellum has three anterior–posterior divisions (Kandel et al., 2000). The primary fissure separates the anterior lobe from the posterior lobe and the posterolateral fissure separates the posterior and flocculo-nodular lobe. Three mediolateral regions that are important functionally are distinguished by two longitudinal grooves that define an elevated ridge in the midline known as the vermis and the cerebellar hemispheres on either side of the vermis. Each of the 26 cerebellar regions distinguished by the AAL template was assigned to one of seven cerebellar anatomical regions of interest (ROI; Table 2), including the anterior and posterior vermis, the bilateral anterior and posterior cerebellar hemispheres and the flocculo-nodular lobe (see Figure 1, no. 3; 6; 1 and 2; 4 and 5; 7 respectively).
Table 1 | Demographic and clinical characteristics.

|                                | Healthy control subjects (N = 41) | Siblings of patients (N = 67) | Schizophrenia patients (N = 62) |
|--------------------------------|----------------------------------|-------------------------------|--------------------------------|
| Age, mean (SD)                 | 30.4 (8.8)                       | 29.8 (8.0)                    | 31.2 (5.8)                     |
| Gender, M/F                    | 19/22                            | 34/33                         | 52/10*                         |
| Handedness*, R/L               | 34/7                             | 60/7                          | 54/5                           |
| Highest degree of educationb  | 5.5 (1.4)                        | 5.2 (2.1)                     | 4.1 (2.1)*                     |
| IQc (SD)                       | 110.1 (15.4)*                    | 102.7 (14.9)*                 | 93.7 (13.5)*                   |
| Duration of illness, years (SD)|                                  |                               | 7.5 (4.3)                      |
| Diagnosis                      |                                  |                               |                                |
| Schizophrenia, N (%)           | 44 (71.0)                        |                               |                                |
| Schizotypiform disorder, N (%) | 3 (4.8)                          |                               |                                |
| Schizoaffective disorder, N (%)| 8 (12.9)                         |                               |                                |
| Otherd, N (%)                  | 7 (11.3)                         |                               |                                |
| Schizotypal features, mean (SD)range | 0.19 (0.18) [0–0.8]           | 0.17 (0.15) [0–0.5]           |                                |
| PANSS symptomsf                |                                  |                               |                                |
| Positive, mean (SD) [range]    | 10.8 (4.0) [7–24]                |                               |                                |
| Negative, mean (SD) [range]    | 11.8 (3.8) [7–23]                |                               |                                |
| Total, mean (SD) [range]       | 61.8 (16.3) [41–108]             |                               |                                |
| Antipsychotic medicationf      |                                  |                               |                                |
| Atypical, N                    | 43                               |                               |                                |
| Risperidone, N; HEQ dose (SD)  | 11; 5.8 (2.5)                    |                               |                                |
| Olanzapine, N; HEQ dose (SD)   | 17; 7.8 (4.2)                    |                               |                                |
| Quetiapine, N; HEQ dose (SD)   | 5; 19.2 (16.6)                   |                               |                                |
| Clozapine, N; HEQ dose (SD)    | 8; 13.7 (6.1)                    |                               |                                |
| Aripiprazole, N; HEQ dose (SD) | 2; 9.0 (4.2)                     |                               |                                |
| Typical, N                    | 5                                |                               |                                |
| Haloperidol, N; HEQ dose (SD)  | 2; 3.5 (0.7)                     |                               |                                |
| Other typicalg, N; HEQ dose (SD)| 3; 4.7 (4.0)                    |                               |                                |
| No current antipsychotic therapy, N | 8                              |                               |                                |

*a Data missing for three patients; b Ranging from no education (0) to university (8). c Estimated intelligence quotient (IQ). d Other diagnoses include delusional disorder (N = 2), brief psychotic disorder (N = 2), and psychosis not otherwise specified (N = 3). e Positive and negative syndrome scale (PANSS), data missing for three patients; f Average haloperidol equivalent (HEQ) dose (mg); g other typical medication includes flupentixol (N = 1), perhenazine (N = 1), and penfluridol (N = 1), data missing for five patients; * indicates a statistically significant difference (at p < 0.05).

Table 2 | Automated anatomical labeling regions per cerebellar ROI.

| L anterior hemisphere | R anterior hemisphere | Anterior vermis | L posterior hemisphere | R posterior hemisphere | Posterior vermis | Flocculo-nodular lobe |
|-----------------------|-----------------------|-----------------|------------------------|------------------------|-----------------|------------------------|
| L lobule III of cerebellar hemisphere | R lobule III of cerebellar hemisphere | Lobule I, II of vermis | L lobule VI of cerebellar hemisphere | R lobule VI of cerebellar hemisphere | Lobule VI of vermis | L lobule X of cerebellar hemisphere (floculus) |
| L lobule IV, V of cerebellar hemisphere | R lobule IV, V of cerebellar hemisphere | Lobule III of vermis | L crus I of cerebellar hemisphere | R crus I of cerebellar hemisphere | Lobule VII of vermis | R lobule X of cerebellar hemisphere (floculus) |
|                        |                       | Lobule IV, V of vermis | L crus II of cerebellar hemisphere | R crus II of cerebellar hemisphere | Lobule VIII of vermis | Lobule X of vermis (nodosus) |

The 26 cerebellar regions distinguished by the automated anatomical labeling (AAL) atlas were assigned to one of seven cerebellar regions of interest; L, left; R, right.
computed for each cerebellar ROI. Next, weighted correlation maps per subject group were transforms were not applied as the correlations were normally dis-
series may lead signals to phase out when averaged. Fisher $r$-$Z$
than one functional region and averaging distinct regional time-
sequently averaged for each ROI (rather than averaging at the

**FUNCTIONAL CONNECTIVITY ANALYSES**

Figure 2 illustrates the consecutive steps of the performed func-
tional connectivity analysis. For each subject, 116 regional mean
time-series were computed by averaging the voxel-based time-
series within each of the anatomically defined regions. Next,
interregional correlation in resting-state time-series between each
possible pair of the 26 cerebellar regions and the 90 cerebral
regions in the AAL template was computed (Figure 2A). For each
subject, functional connectivity per cerebellar ROI was then calcu-
lated by averaging the correlation coefficients of the AAL regions
within each of the seven cerebellar regions (Figure 2B), render-
ing weighted correlation coefficients $r_{ij}$ for the connections $[i,$
$j]$ of each cerebellar ROI $i$ ($N = 7$) with each cerebral region $j$
($N = 90$). Interregional correlations were first computed and sub-
sequently averaged for each ROI (rather than averaging at the
level of regional time-series) as the ROIs may comprise more
than one functional region and averaging distinct regional time-
series may lead signals to phase out when averaged. Fisher $r$-$Z$
transforms were not applied as the correlations were normally dis-
tributed. Next, weighted correlation maps per subject group were
computed for each cerebellar ROI (Figure 2C), rendering three
matrices (for control, sibling, and patient groups) per ROI, with
all 90 connections $[i, j]$ of the particular ROI $i$ on the $x$-axis and
the number of subjects in the group ($N = 41; 67; 62$ respectively)
on the $y$-axis.

**HEALTHY CEREBELLAR FUNCTIONAL CONNECTIVITY**

In order to interpret possible differences in cerebellar functional
connectivity between subject groups, the general pattern of func-
tional connections per cerebellar ROI was investigated in the
healthy control subjects. As functionally connected regions have
been shown to exhibit a high degree of temporal coherence (Biswal
et al., 1995; Van den Heuvel et al., 2008), higher interregional
correlation in time-series was interpreted as a higher degree of
functional connectivity between two regions.

**CEREBELLAR FUNCTIONAL CONNECTIVITY IN SCHIZOPHRENIA PATIENTS AND SIBLINGS**

Differences between subject groups in cerebellar functional con-
nectivity patterns were examined using two functional connectiv-
ity measures. First, overall connectivity strength per cerebellar ROI
was computed, providing information on the global level of com-
unication between each cerebellar ROI and the rest of the brain.

**Analysis 1 overall connectivity strength $S$ per cerebellar ROI**

Connectivity strength $S_i$ of each cerebellar ROI $i$ was computed
as the average of all correlations between region $i$ and all extra-
cerebellar regions $j$, providing information on the total level of
connectivity of each cerebellar ROI (Figure 2D). Formally:

$$ S_i = \frac{\sum F_{ci,j}}{N_j} $$

Overall connectivity strength is a global measure of the extent
to which the ROI is integrated in the brain network. However, it is
not specific as to whether any particular connections are affected.
Therefore, a pairwise approach was used next, to “zoom in” on
discrete connections.

**Analysis 2 functional connectivity $F_c$ of discrete connections**

Computed as the (averaged) correlation of each cerebellar region $i$
with each extra-cerebellar region $j$ of the brain network, each func-
tional connection $F_{ci,j}$ is an element of the set $F_C$ of all functional
connections ($N = 90$) of the cerebellar ROI (Figure 2E):

$$ F_{ci,j} \in F_C $$

In both analyses, the actual values (i.e., both positive and neg-
ative correlations) were used when averaging across correlation
coefficients. As a consequence, negative correlations – which were
both scarce and small in amplitude (all mean correlation coeffi-
cients $> -0.1$) – were interpreted as lower levels of functional
connectivity.

To exclude potential bias (in variance estimates) of family rela-
tionships within and between subject groups, the analyses were
repeated with only unrelated individuals ($N = 132$), i.e., 46 schiz-
ophrrenia patients, 47 siblings of patients, and 39 healthy control
subjects.

**CLINICAL CORRELATES OF CEREBELLAR CONNECTIVITY**

Using linear regression analyses, the association between any possi-
ble changes in cerebellar connectivity measures and clinical vari-
ables (i.e., duration of illness; severity of PANSS symptoms and
IQ) was investigated, and the dose of antipsychotic medication (in
haloperidol equivalent) at the time of scanning was examined as a
potential confounder.

**STATISTICAL ANALYSIS OF GROUP-DIFFERENCES**

Connectivity strength $S$ of each cerebellar ROI and functional
connectivity $F_c$ of each individual connection of schizophrenia
patients and siblings of schizophrenia patients were compared to those of the healthy comparison subjects. To examine the statistical significance of group-differences, permutation testing was used (5000 permutations; Bassett et al., 2008; Lynall et al., 2010; Van den Heuvel et al., 2010). To this end, random permutation of group assignment was performed, maintaining the original number of subjects per group, rendering three groups of 41, 62, and 67 randomly assigned subjects. Consequently, weighted correlation maps per randomly assigned subject group were computed for each cerebellar ROI. Using the resulting correlation maps, the connectivity measures were recalculated. This process was repeated 5000 times, resulting in a between-group difference null distribution of the connectivity measures. Finally, the observed differences between the original subject groups were compared to the normal distribution of differences after random permutation, to explore the null hypothesis that the observed differences were not determined by subject group membership. Finally, $p$-values were assigned to the group effects by computing (after random permutation) the percentage of findings that was more extreme than the observed difference between the original subject groups. In view of the number of tests performed in the connectivity strength analysis (Analysis 1, Figure 2D), a $p$-value of $<0.01$ was considered to

**FIGURE 2 | Functional connectivity analysis.** Consecutive steps of the functional connectivity analysis. (A) Computation of correlation between cerebellar and cerebral resting-state fMRI time-series, (B) calculation of correlation coefficients of cerebellar ROIs, (C) computation of weighted correlation maps per subject group and per cerebellar ROI. Next, computation of (D) overall connectivity strength per cerebellar ROI, and (E) functional connectivity of discrete connections.
RESULTS

HEALTHY CEREBELLAR FUNCTIONAL CONNECTIVITY

In the healthy comparison subjects, mean functional connectivity of the posterior hemispheres [mean Fc (SD) = 0.27 (0.07)] > anterior hemispheres [mean Fc (SD) = 0.21 (0.06)] and the right cerebellum [mean Fc (SD) = 0.27 (0.08)] > left cerebellum [mean Fc (SD) = 0.22 (0.06)] (both p < 0.0001). Furthermore, mean functional connectivity of the lateral cerebellum (i.e., hemispheres) [mean Fc (SD) = 0.24 (0.06)] was significantly higher than that of midline structures (i.e., vermis and flocculo-nodular lobe) [mean Fc (SD) = 0.14 (0.05)] (p < 0.0001).

The highest levels of functional connectivity were found between the right anterior cerebellar hemisphere and right lingual gyrus (mean Fc = 0.54, SD 0.14), hippocampus [mean Fc (SD) = 0.50 (0.18)] and parahippocampal gyrus [mean Fc (SD) = 0.50 (0.15)]. Overall, cerebellar connections to posterior medial regions of the cerebral cortex [i.e., (pre)cuneus, calcarine sulcus, lingual gyrus] showed high levels of functional connectivity, irrespective of cerebellar ROI [mean Fc (SD) of all ROIs 0.26 (0.08), 0.26 (0.10); 0.34 (0.08), 0.32 (0.08); 0.32 (0.09), 0.29 (0.10); 0.29 (0.09), 0.33 (0.12); for the left and right cuneus; precuneus; calcarine sulcus; and lingual gyrus respectively]. The lowest levels of functional connectivity were observed for connections with the bilateral lentiform nucleus [mean Fc (SD) = 0.11 (0.15)], in particular the globus pallidus [left; right mean Fc (SD) = 0.05 (0.15); 0.09 (0.15)], and with the gyrus rectus [left; right mean FC (SD) = 0.11 (0.17); 0.09 (0.17)] (Figure 3).

ANALYSIS 1 OVERALL CONNECTIVITY STRENGTH S OF CEREBELLAR ROIs

Overall connectivity strength of the posterior and flocculo-nodular lobes of the cerebellum with the rest of the brain did not differ between groups, suggesting intact functional connectivity of posterior and flocculo-nodular cerebellar regions in schizophrenia. Connectivity strength of the right anterior hemisphere of the cerebellum however, was found to be 23% less (p = 0.01) in schizophrenia patients compared to healthy comparison subjects. Furthermore, on trend level, connectivity strength of the anterior vermis was less in both schizophrenia patients (−27%, p = 0.04) and siblings (−22%, p = 0.05), compared to healthy comparison subjects (Figure 4).
ANALYSIS 2 FUNCTIONAL CONNECTIVITY OF DISCRETE CONNECTIONS: PATIENTS VERSUS CONTROLS

Impaired functional connectivity was most pronounced in the connections between the right anterior cerebellar hemisphere and vermis, and left-cerebral regions. Particularly between the right anterior vermis and the left-sided triangular part of the inferior frontal gyrus ($F_c = -0.13, p = 0.0024$), supplementary motor area ($F_c = -0.12, p = 0.0014$), middle cingulate gyrus ($F_c = -0.12, p = 0.0004$), hippocampus ($F_c = -0.17, p = 0.0004$), and thalamus ($F_c = -0.16, p < 0.0001$). Functional connectivity between the latter two regions (i.e., left hippocampus and thalamus) and the anterior cerebellar vermis was also decreased in patients ($F_c = -0.14, p < 0.0001; F_c = -0.13, p = 0.0008$ respectively). The only significant increase in functional connectivity in schizophrenia patients relative to controls was found between the posterior vermis and left lingual gyrus ($F_c = 0.15, p < 0.0001$). All these findings survived FDR-correction (Figure 5).

ANALYSIS 2 FUNCTIONAL CONNECTIVITY OF DISCRETE CONNECTIONS: SIBLINGS VERSUS CONTROLS

Compared to schizophrenia patients, the siblings demonstrated both similarities and disparities in functional connectivity of the...
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cerebellum. Siblings showed decreased functional connectivity (compared to control subjects) between the anterior cerebellar vermis and the triangular part of the left inferior frontal gyrus ($F_c = -0.08, p = 0.009$), left insula ($F_c = -0.08, p = 0.007$), and left hippocampus ($F_c = -0.11, p = 0.002$), as well as between the right anterior cerebellar hemisphere and left hippocampus ($F_c = -0.12, p = 0.003$), resembling the findings in patients. However, these findings were not as strong as in patients, as they failed to reach whole-brain significance (note that impaired functional connectivity between the anterior cerebellar vermis and left insula did not reach whole-brain significance in both siblings and patients), and should be considered exploratory. Conversely, functional connectivity to other brain regions that were found to be disconnected from the cerebellum in schizophrenia patients (e.g., left thalamus, middle cingulate gyrus, and supplementary motor area) was not decreased in siblings of schizophrenia patients, compared to the healthy control subjects. Table 3 depicts all findings at $p < 0.01$ in schizophrenia patients and siblings of patients.

Importantly, as siblings were psychiatrically healthy (see inclusion criteria and SIS-R scores, Table 1) and, on average, around 30 years of age, it is unlikely that any would still convert to illness. Lastly, repeating the functional connectivity analyses in a reduced

| Table 3 | Absolute differences in functional connectivity between subject groups. |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| **Cerebellar regions** | **Sz** | **Sib** | **Sz** | **Sib** | **Sz** | **Sib** | **Sz** | **Sib** |
| **CEREBRAL REGIONS** | **L anterior hemisphere** | | **R anterior hemisphere** | | **Anterior vermis** | | **L posterior hemisphere** | | **R posterior vermis** | | **Posterior lobe** | | **Flocculo-nodular lobe** | |
| R superior frontal gyrus, dorsolateral | 0.10 | | | | | | | | | | | | |
| L superior frontal gyrus, orbital | 0.11 | 0.07 | | | | | | | | | | | |
| L middle frontal gyrus, orbital | 0.10 | | | | | | | | | | | | |
| L inferior frontal gyrus, triangular | 0.13* | 0.09 | 0.08 | | | | | | | | | | |
| L supplementary motor area | 0.12* | 0.09 | | | | | | | | | | | |
| R supplementary motor area | 0.10 | | | | | | | | | | | | |
| R superior frontal gyrus, medial | 0.09 | | | | | | | | | | | | |
| L superior frontal gyrus, medial/orbital | 0.08 | | | | | | | | | | | | |
| L gyrus rectus | 0.09 | | | | | | | | | | | | |
| R gyrus rectus | | 0.07 | | | | | | | | | | | |
| L insula | | 0.10 | 0.08 | | | | | | | | | | |
| L anterior cingulate gyrus | | | 0.12* | 0.10 | | | | | | | | | |
| L middle cingulate gyrus | | | 0.12* | 0.10 | | | | | | | | | |
| L hippocampus | | | 0.17* | 0.12 | 0.14* | 0.11 | | | | | | 0.11 |
| L lingual gyrus | | | | 0.15* | 0.08 | | | | | | | |
| R lingual gyrus | | | | | 0.10 | | | | | | | | |
| R inferior occipital gyrus | | 0.11 | | | | | | | | | | | |
| L inferior parietal lobe | | 0.10 | | | | | | | | | | | |
| L paracentral lobe | | | 0.09 | | | | | | | | | | |
| L caudate nucleus | | | | 0.08 | | | | | | | | | |
| R caudate nucleus | | | | 0.09 | | | | | | | | | |
| R globus pallidus | | 0.10 | | | | | | | | | | | |
| L thalamus | | 0.16* | 0.13* | | 0.11 | | | | | | | | |
| L superior temporal gyrus | | 0.11 | | | | | | | | | | | |
| R superior temporal pole | | 0.11 | | | | | | | | | | | |

Absolute difference in interregional correlation coefficients between schizophrenia patients and healthy comparison subjects; and siblings of patients and healthy comparison subjects. Decreased functional connectivity relative to healthy comparison subjects is displayed in blue, increased functional connectivity in red. Findings at $p < 0.01$ are depicted; * indicates whole-brain significance after FDR-correction. Sz, schizophrenia patients; Sib, siblings of schizophrenia patients, L, left-sided, R, right-sided.
number of study participants that were completely unrelated did not alter the findings.

**CLINICAL CORRELATES OF CEREBELLAR CONNECTIVITY**

Cerebellar functional connectivity measures that were significantly different in schizophrenia patients, compared to healthy comparison subjects, were investigated for associations with medication dose, duration of illness, symptom severity, and global cognitive performance, using linear regression analysis. No association between the dose of antipsychotic medication at the time of scanning and cerebellar functional connectivity was found (all p > 0.420). After multiple comparison correction, there were no significant associations between aberrant cerebellar functional connectivity measures and clinical variables. On trend level, a counter-intuitive association between more negative symptoms and increased Fc between the cerebellar right anterior hemisphere and left hippocampus was observed (**Table 4**).

**DISCUSSION**

The main finding of our study is the presence of an aberrant level of functional connectivity of the cerebellum in schizophrenia patients and their healthy siblings. Our findings suggest that in schizophrenia, the cerebellum, specifically the vermis, and right anterior hemisphere, is functionally disconnected from a range of left-cerebral cortical and subcortical regions, including frontal, cingulate, and occipital regions, as well as the thalamus and hippocampus. Importantly, unaffected siblings of schizophrenia patients demonstrated several similarities in cerebellar functional connectivity compared to patients, in particular reduced functional connectivity of the cerebellum to the left hippocampus. This overlap suggests that cerebellar dysconnectivity may be related, at least in part, to familial (and possibly genetic) factors.

Disparities in cerebellar connectivity between schizophrenia patients and their siblings were also found. Of these, the dissimilarity in cerebellar–thalamic dysconnectivity may be the most meaningful, as the thalamus is the obligatory relay for all efferent cerebellar projections to the cortex. In this context, our findings may imply that cerebellar–thalamic dysconnectivity might be related more to the manifestation of the illness, than to familial (or genetic) factors.

Although the term “dysconnectivity” emphasizes the notion of abnormal, rather than necessarily decreased, functional integration between brain regions (Stephan et al., 2009a), structural and functional connectivity studies in schizophrenia have mostly reported reduced (rather than increased) connectivity in schizophrenia (Pettersson-Yeo et al., 2011). In line with these findings, our study shows that in schizophrenia, functional connectivity of the cerebellum is mostly decreased. Furthermore, studies have reported changing patterns of reduced connectivity across the different stages of disease, from chronic schizophrenia to individuals at high (genetic) risk for psychosis (including healthy first-degree relatives of schizophrenia patients; Pettersson-Yeo et al., 2011). These findings support the presently reported overlap in cerebellar dysconnectivity between patients and siblings. Additional support for this finding comes from a recent study examining functional connectivity within and between four predefined functional networks: the default mode network and predefined functional networks: the default mode network and cerebellar (or genetic) factors.

Intriguingly, greater reductions in connectivity between the frontal–parietal and cerebellar regions were found to be robustly predictive of worse cognitive performance across groups and predictive of more disorganization symptoms among patients. The association between cognitive performance and cerebellar functional connectivity is not replicated by the present study. However, as modest cognitive deficits are reported in adult patients with (especially right-sided) cerebellar lesions, while similar lesions in children lead to pervasive impairments (suggesting a critical role for the cerebellum in development; Alexander et al., 2011), the relationship between cerebellar abnormalities and cognition may not be straightforward and may depend on factors such as age of illness onset. Regarding the relationship with clinical symptoms, the meaning of the counter-intuitive trend between greater severity of PANSS

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**Table 4 | Clinical correlates of aberrant functional connectivity measures.**

| Aberrant functional connectivity measures | S Cer2 | S Cer3 | Fc Cer2-IFt | Fc Cer2-SMA | Fc Cer2-MCin | Fc Cer2-Hip | Fc Cer2-Tha | Fc Cer3-Hip | Fc Cer3-Tha | Fc Cer6-Lin |
|---|---|---|---|---|---|---|---|---|---|---|
| Duration of illness | −0.21 | −0.07 | −0.29 | −0.26 | −0.15 | −0.05 | −0.10 | 0.10 | 0.12 | −0.22 |
| PANSS Total score | 0.24 | 0.12 | 0.09 | 0.02 | 0.29 | 0.33 | 0.31 | 0.26 | 0.10 | −0.10 |
| Positive symptoms | 0.16 | 0.15 | −0.01 | 0.01 | 0.23 | 0.17 | 0.26 | 0.16 | 0.03 | −0.01 |
| Negative symptoms | 0.17 | 0.07 | 0.05 | −0.01 | 0.22 | 0.37* | 0.27 | 0.25 | 0.18 | −0.07 |
| IQ | −0.02 | −0.04 | −0.15 | −0.02 | −0.03 | −0.29 | −0.16 | −0.11 | −0.07 | −0.12 |

*Standardized correlation coefficients (ß) describing the correlation between clinical variables and cerebellar functional connectivity metrics that were found to be aberrant in schizophrenia patients compared to healthy comparison subjects (**p < 0.01, uncorrected**). S = connectivity strength; Fc = functional connectivity; Cerebellar ROI 2 (Cer2) = right anterior cerebellar hemisphere; Cerebellar ROI 3 (Cer3) = Anterior vermis; IFt, inferior frontal gyrus, triangular part; SMA, supplementary motor area, MCin, middle cingulate gyrus; Hip, hippocampus, Tha, thalamus, Lin, lingual gyrus.*
negative symptoms and increased functional connectivity between the cerebellar anterior right hemisphere and left hippocampus is unclear, but may be an indication of discrete symptom dimensions in schizophrenia with distinct underlying neurobiology (Ke et al., 2010).

Our findings suggest that functional dysconnectivity of the cerebellum to the cerebrum involves mainly anterior and vermal areas of the cerebellum. The vermis has previously been labeled the “limbic cerebellum,” as the (anterior) vermis is the principal cerebellar target of limbic projections (Schmahmann, 2000). Furthermore, behavioral studies support a relationship between cerebellar midline structures and the modulation of emotion (Heath and Harper, 1974; Stoodley and Schmahmann, 2010), and lesions of the vermis have been shown to produce affective symptoms, ranging from emotional blunting and depression to disinhibition and psychotic features (Schmahmann and Sherman, 1998). The observed reduction in functional integration between this cerebellar region and limbic regions such as the hippocampus, cingulate cortex and (anterior nuclei of) the thalamus, could perhaps be interpreted in this context. Notably, the present study found no evidence for impaired functional connectivity of the posterior cerebellar hemispheres, while these cerebellar regions have been preferentially linked to cognitive processing (Habas et al., 2009; Krienen and Buckner, 2009). Cerebellar involvement in schizophrenia may be either limited to the vermis and anterior lobe, or involvement of the posterior cerebellar hemispheres in (cognitive deficits in) schizophrenia may be more subtle, attenuating in our whole-brain approach with corresponding stringent correction for multiple comparisons. Alternatively, as a distinct topographic organization of the cerebellum has been proposed in which certain areas of the cerebellar cortex interact specifically with certain areas of the cerebral cortex (Stoodley and Schmahmann, 2010; Buckner et al., 2011), the relatively large posterior hemispheres (compared to the other cerebellar ROIs) may have such distributed functional connections, that “picking up” on dysconnectivity of any particular connection(s) was precluded by our definition of ROIs.

The neuronal basis for dysconnectivity remains to be established. It could result from either aberrant wiring of connections during development or from impaired synaptic plasticity (or both; Stephan et al., 2009a). As studies have suggested a link between functional and structural brain connectivity (Hagmann et al., 2008; Honey et al., 2009; Van den Heuvel et al., 2009), our findings may reflect impaired structural connections (i.e., white matter tracts) between affected brain regions. Indeed, reduced fractional anisotropy (FA), commonly interpreted as reduced integrity, of cerebellar white matter tracts has been shown in schizophrenia (Kanaan et al., 2009; Kyriakopoulos and Frangou, 2009). Accompanied by normal mean diffusivity, this FA reduction is most likely due to disordered microstructural architecture, rather than disordered myelination (Kanaan et al., 2009). Accordingly, one study using magnetic resonance spectroscopic imaging reported decreased levels of a putative neuronal/axonal marker in the anterior cerebellar vermis, suggesting dysfunction or loss of neurons in that region (Deicken et al., 2001). Cerebellar dysconnectivity may thus result from decreased neurons or disordered neuronal architecture in the cerebellum.

Some issues have to be taken into account when interpreting the results of our study. First, medicated patients were studied and antipsychotic mediation has been shown to affect cerebellar functional connectivity (Stephan et al., 2009b). However, the haloperidol equivalent dose of antipsychotic medication at the time of scanning was not associated with functional connectivity measures and the unmedicated siblings of schizophrenia patients showed similar abnormalities in cerebellar functional connectivity, suggesting that cerebellar dysconnectivity is unlikely to be due to antipsychotic medication alone. Second, there was a preponderance of men in the schizophrenia patients, which was not paralleled in the healthy control and sibling groups. Nonetheless, as we applied permutation testing, it is very unlikely that our findings were driven by any other factor than group membership (i.e., patient, sibling, or control), as group-differences determined by other factors – such as gender – would not have been among the most extreme findings after random permutation of group assignment and thus not be deemed significant. Furthermore, our subdivision of the cerebellum may not be the optimal division of the cerebellum in terms of its functional connections. As the connective properties of the cerebellum (i.e., connected to the cerebral cortex only by ways of polysynaptic connections) are relatively unamenable to traditional anatomical methods, the connectional topography of the cerebellum remains largely unmapped (O’Reilly et al., 2010). Recent studies using transneuronal tracing techniques and functional neuroimaging have provided some insight into the functional organization of the cerebellar cortex, but both distinct and overlapping functional zones of the cerebellar cortex have been reported and findings diverge between studies (O’Reilly et al., 2010; Stoodley and Schmahmann, 2010; Buckner et al., 2011). It was therefore decided to adhere to a broad division of the cerebellum, based on gross anatomy. Moreover, although various preprocessing steps were used to deal with potential confounds associated with resting-state fMRI (e.g., cardiorespiratory oscillations, head movement, scanner noise), influences of these factors cannot be ruled out. There is also an inherent risk of potential bias associated with cortical parcellation methods such as the AAL template, as the spatial scale of nodal parcellation has been shown to influence brain network properties (Wang et al., 2009; Fornito et al., 2010). New approaches are being developed to perform whole-brain connectivity mapping using individual pairs of voxels, without the need of arbitrary parcellation of the cortex (Zalesky et al., 2010), but interpreting voxels as distinct information processing units has its own limitations and inevitably, investigations of in vivo whole-brain functional connectivity have a resolution limit (Wig et al., 2011). Finally, although we interpret our findings of cerebellar dysconnectivity in schizophrenia in terms of underlying neurobiology and psychopathology, it should be noted that cause and effect cannot be derived from our cross-sectional data.

In conclusion, our study indicates that the cerebellum, in particular the vermis and right anterior hemisphere, is functionally dysconnected in schizophrenia. Furthermore, the observed overlap in cerebellar dysconnectivity between schizophrenia patients and their healthy siblings suggest that cerebellar dysconnectivity is related, at least in part, to familial risk for psychosis. Whether, and if so to what extent, cerebellar dysconnectivity is mediated by genetic factors needs to be established in genetic studies.
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