Progressive Loss of Cerebellar Volume in Childhood-Onset Schizophrenia

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Objective: Childhood-onset schizophrenia is a severe and unremitting form of the disorder. Prospective brain magnetic resonance imaging (MRI) studies have found progressive loss of total cerebral volume during adolescence, primarily attributable to accelerated loss of cortical gray matter. Because there is evidence of cerebellar involvement in schizophrenia, the authors examined cerebellar volume and its relation to cortical gray matter development during adolescence in patients with childhood-onset schizophrenia and healthy comparison subjects.

Method: Total cerebellar volume was algorithmically calculated for 108 anatomical brain MRI scans from 50 patients (20 of whom were female) and 101 scans from 50 age- and gender-matched healthy volunteers (20 of whom were female). The age range of the patients and comparison subjects was 8 to 24. Midsagittal vermal area and posterior-inferior vermal lobe volume were measured by hand. Prospective rescans were obtained at approximately 2-year intervals. Cross-sectional and longitudinal data were combined in mixed model regressions to compare developmental changes for the groups.

Results: In contrast to healthy volunteers, patients with schizophrenia showed a progressive loss of cerebellar volume during adolescence. Cerebellar and cerebral volume decreases were significantly correlated in childhood-onset schizophrenia.

Conclusions: Childhood-onset schizophrenia is associated with significant progressive loss of cerebellar volume during adolescence, consistent with previously reported decreases in total cerebral and cortical gray matter. At least in these patients with severe early-onset schizophrenia, the loss appears secondary to a generalized process.
ment in cerebellar and posterior-inferior vermal volumes would be seen throughout adolescence. We were also interested in whether cerebellar changes paralleled previously reported cortical gray matter loss for this group.

Method

Childhood-Onset Schizophrenia

Children and adolescents were recruited nationally for an ongoing study of childhood-onset schizophrenia (34). Inclusion criteria were a DSM-III-R or DSM-IV diagnosis of schizophrenia with onset of psychosis before age 12, a premorbid full-scale IQ of at least 70, the absence of active medical or neurological disease, and poor response to or inability to tolerate treatment with at least two different neuroleptics. The diagnosis was established by using previous records and clinical and structured interviews of the children and parents as described in detail elsewhere (35).

The study group of 50 patients with childhood-onset schizophrenia for whom at least one scan was available included 20 females. At baseline and at each follow-up visit, subjects were administered the WISC-R, the WISC-III, or the WAIS-R as appropriate. At follow-up, WISC raw scores for information and comprehension subscales were obtained to compare performance on these scales independent of subject age. A previously described method (2) was used to review premorbid neuropsychological, school, and medical records. Twenty-seven patients (nine of whom were female) had developmental language difficulties, and nine had comorbid pervasive developmental disorder, not otherwise specified, according to the Autism Screening Questionnaire (36).

Analyses of initial scans included 50 scans per diagnostic group. Patients returned for rescan approximately every 2 years. In the childhood-onset schizophrenia group, 19 patients had two, 12 had three, and five had four scans. In the comparison group, 19 subjects had two, 13 had three, and two had four scans. Developmental analyses were based on 108 childhood-onset schizophrenia scans and 101 comparison group scans. Using only subjects with more than one scan, we examined the relationship between cerebellar volume change and clinical or anatomical measures for 34 healthy volunteers and 36 patients with childhood-onset schizophrenia. All patients with childhood-onset schizophrenia were taking neuroleptics at follow-up, most frequently a combination of clozapine and at least one other antipsychotic medication (N=14), clozapine alone (N=12), or antipsychotic drugs other than clozapine (N=10).

Healthy Volunteers

Fifty healthy children and adolescents matched for age, sex, and handedness were recruited from the community (Table 1). Structured rating scales and interviews of child and parents were performed as described elsewhere (37).

Assent from the child and written consent from the parents were obtained for both patients and comparison subjects. The NIMH Institutional Review Board approved this study.

MRI Image Acquisition

All subjects were scanned on the same GE 1.5-T Signa scanner (GE Medical Systems, Milwaukee). T1-weighted images with contiguous 1.5-mm slices in the axial plane and 2.0-mm slices in the coronal plane were obtained by using three-dimensional spoiled gradient recalled echo in the steady state. Imaging parameters were echo time=5 msec, repetition time=24 msec, flip angle=45°, acquisition matrix=192×256, number of excitations=1, and field of view=24 cm. Head placement was standardized as previously described (3).

<table>
<thead>
<tr>
<th>TABLE 1. Characteristics at Initial MRI Scan and Rescan Data for 50 Patients With Childhood-Onset Schizophrenia and 50 Healthy Comparison Subjects</th>
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</thead>
<tbody>
<tr>
<td>Characteristic</td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>Age (years)</td>
</tr>
<tr>
<td>Age at onset of schizophrenia (years)</td>
</tr>
<tr>
<td>Height (cm)a</td>
</tr>
<tr>
<td>Weight (kg)b</td>
</tr>
<tr>
<td>WISC standard scores</td>
</tr>
<tr>
<td>Block designc</td>
</tr>
<tr>
<td>Previous neuroleptic exposure (months)b</td>
</tr>
<tr>
<td>Hospitalization (months)</td>
</tr>
<tr>
<td>Number of scans</td>
</tr>
<tr>
<td>Interval between scans (years)</td>
</tr>
<tr>
<td>Gender</td>
</tr>
<tr>
<td>Male</td>
</tr>
<tr>
<td>Female</td>
</tr>
<tr>
<td>Handedness</td>
</tr>
<tr>
<td>Right</td>
</tr>
<tr>
<td>Left</td>
</tr>
<tr>
<td>Not known</td>
</tr>
<tr>
<td>Comorbid pervasive developmental disorder</td>
</tr>
</tbody>
</table>

a Significant difference between groups (t=–2.07, df=92, p=0.04).
b Data available for 45 patients.
c Data available for 49 patients.
d Significant difference between groups (t=–8.51, df=87, p=0.0001).
d Significant difference between groups (t=–8.63, df=90, p=0.0001).

Image Analysis

Cerebrum and cerebellum. Total cerebral and cerebellar volumes were quantified by using a three-part fully automated image analysis process described in detail elsewhere (33, 38). Reliability of the algorithms equaled unity for test-retest with the same scan. Intraclass correlation coefficients (ICC) for 10 subjects rescanned after leaving and reentering the magnet exceeded 0.98 for both measures. Automated and hand-traced measures of cerebellar volume were highly reliable (N=17, ICC=0.94).

Vermis. Midsagittal vermal area and posterior-inferior vermal lobe volume, excluding the cerebellar tonsils, were hand-measured by one rater (A.C.V.) using the MNI Display program (Montreal Neurological Institute, McGill University, Montreal) on a Silicon Graphics workstation (Mountain View, Calif.). Beginning with the cerebral midsagittal slice, the best vermis midsagittal slice, which differed from the midcerebellar 63% of the time (39), was selected in sagittal view and confirmed in coronal and axial views (Figure 1). Areas for anterior, superior-inferior, and posterior-inferior lobes were hand-traced on the vermis midsagittal slice and added to derive total vermis area. Boundaries for the anterior lobe are the primary fissure and fourth ventricle; boundaries for the superior-inferior lobe were the primary and prepyramidal fissures; boundaries for the posterior-inferior lobe (lobules VIII–X) were the prepyramidal fissure and fourth ventricle (31, 40).

Posterior-inferior volume was obtained by starting with the midsagittal outline that provided superior and inferior landmarks on each coronal slice. Lateral boundaries for each coronal slice.
The best vermis midsagittal slice was selected in sagittal view and confirmed in coronal and axial views with the aid of the crosshair cursor. Posterior-inferior volume was obtained by starting with the midsagittal outline that provided superior and inferior landmarks on each coronal slice. Lateral boundaries for each coronal slice were the interface between CSF and gray matter, and filled coronal areas were checked slice by slice in axial and sagittal views.

**TABLE 2. MRI Regional Anatomical Volumes Based on Initial Scans of 50 Patients With Childhood-Onset Schizophrenia and 50 Healthy Comparison Subjects**

<table>
<thead>
<tr>
<th>Brain Region Measure</th>
<th>Patients With Childhood-Onset Schizophrenia</th>
<th>Comparison Subjects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total cerebral volume (cm³)</td>
<td>1068.72 ± 121.00</td>
<td>1099.65 ± 128.85</td>
</tr>
<tr>
<td>Cerbellar volume (cm³)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>131.31 ± 14.75</td>
<td>133.96 ± 12.45</td>
</tr>
<tr>
<td>Right</td>
<td>67.60 ± 7.70</td>
<td>68.77 ± 6.34</td>
</tr>
<tr>
<td>Left</td>
<td>63.70 ± 7.35</td>
<td>65.19 ± 6.38</td>
</tr>
<tr>
<td>Vermal area (cm³)</td>
<td>9.54 ± 1.13</td>
<td>9.89 ± 1.03</td>
</tr>
<tr>
<td>Posterior-inferior vermal area (cm³)</td>
<td>2.39 ± 0.47</td>
<td>2.50 ± 0.46</td>
</tr>
</tbody>
</table>

*Measures included vermal area. No comparisons reached statistical significance.*

Each group included 30 male and 20 female subjects.

...were the CSF-gray matter interface (Figure 1), and filled coronal areas were checked slice by slice in axial and sagittal views. Measurement of volumes of anterior and superior-anterior vermis was not attempted because these regions do not have true lateral boundaries.

Interrater reliability between primary (A.C.V.) and secondary (A.K.) raters was obtained twice to check for rater drift with at least 10 brains per run; mean ICC=0.94 (SD=0.06) for posterior-inferior volume, and mean ICC=0.93 (SD=0.06) for vermal area. Interrater reliability was verified on three occasions: mean ICC=0.87 (SD=0.07) for posterior-inferior volume, and mean ICC=0.93 (SD=0.02) for vermal area.

**Statistical Analysis**

Fisher’s exact and Mann-Whitney test procedures were used to evaluate the comparability of patient and comparison groups.

Polynomial growth models were used to examine growth patterns of brain structures for initial (cross-sectional) scans. The initial cubic model was size=intercept + beta₁* (age–mean age) + beta₂* (age–mean age)² + beta₃* (age–mean age)³ + ε. The model parameters (intercept and beta coefficients) were initially allowed to vary by sex and diagnostic group. The full cubic model was compared with simpler quadratic, linear, and constant models. Once the order of the model was established, testing was performed to determine whether an additive model could replace the interactions between sex and diagnostic group for the height and shape parameters of the curves. Hypothesis tests and model selection were initially based on F statistics. For F statistics with p values less than 0.10 that related to group or sex differences, permutation tests were performed to lessen the likelihood that a significant finding was due to a small number of outliers. To implement this procedure, 500 analyses using the preferred model were performed in which the sex and diagnostic group designation were randomly reassigned. The original F statistic based on the correct group and sex designation was compared with the resulting 500 F statistics, and an empirical p value was obtained.

The same polynomial growth models were used for the analysis of longitudinal data including all of each individual’s scans (41, 42). To account for within-person correlation, intercepts were treated as normally distributed random effects that varied by individual, while beta coefficients for age, age-squared, and age-cubed terms were modeled as fixed effects. Although the statistical information provided by individuals with only one scan may be less than that obtained from those with multiple scans, single scans do provide additional information about between-person variation and overall curve shape.

Spearman correlation was used to examine the relationship between slopes for cerebellar and total cerebral volume loss. Slopes were calculated only for subjects with more than one scan; for those with more than two scans, first and last scans were selected. Stepwise regression and analysis of variance were used to examine clinical and treatment variables in relation to cerebellar volume decrease in childhood-onset schizophrenia.

**Results**

**Initial Scans**

As expected, all volumes were greater for males than females (p<0.02) (Table 2 and Table 3). In analyses limited to each subject’s initial scan, no significant diagnostic differences (Table 2) or sex-by-diagnosis interactions were seen for childhood-onset schizophrenia and healthy comparison subjects for total cerebral volume, cerebellar measures, or vermal measures.

**Developmental Trajectories**

**Total cerebellar volume.** In contrast, as seen in Table 3 and Figure 2, developmental trajectories differed significantly for the childhood-onset schizophrenia group (F=9.30, df=2, 105, p=0.006), with total cerebellar volume de-
creasing with increasing age for patients with childhood-onset schizophrenia but not for comparison subjects. For comparison purposes, and in agreement with previous reports (3), developmental trajectories for total cerebral volume also differed significantly for the childhood-onset schizophrenia group \( (F=16.57, \text{df}=1, 107, p<0.002) \). As seen in Figure 2, total cerebral volume decreased with age for patients with childhood-onset schizophrenia but not for healthy comparison subjects. By visual examination, losses of cerebellar and cerebral volumes appeared to start at approximately the same time.

**Vermal area and posterior-inferior vermal volume.**

No age-related changes were found for either comparison subjects or patients with childhood-onset schizophrenia.

**Cerebellar volume change and anatomical and clinical measures.** Using only childhood-onset schizophrenia subjects with more than one scan, we found that rates \( (\text{cm}^3/\text{year}) \) of decrease of total cerebellar and cerebral volumes were significantly correlated \( (r=0.41, N=36, p=0.01) \). Within the patient group, no significant relationships were observed between cerebellar volume decrease and scores on WISC vocabulary or block design subscales; age at onset; number of months of hospitalization; history of prepsychotic language disorder; pervasive developmental disorder diagnosis; scores on the Scale for the Assessment of Negative Symptoms, Scale for the Assessment of Positive Symptoms, or Brief Psychiatric Rating Scale; medication group (clozapine alone, clozapine with at least one other antipsychotic drug, other antipsychotic); or slopes for change in full-scale IQ or in information and comprehension subscale raw scores.

**Discussion**

In this prospective study of cerebellar morphology in childhood-onset schizophrenia, a progressive loss of cerebellar volume in childhood-onset schizophrenia was found during adolescence, paralleling the loss in total cerebral volume previously reported for this group (3). The cerebellum is one of the first brain structures to begin to differentiate and one of the last to achieve maturity (43), reaching peak volume several years later than total cerebral volume (Giedd et al., unpublished data). However, in patients with childhood-onset schizophrenia, cerebellar loss seems to start at about the same age (within 1 year, by visual examination of confidence intervals) as loss of cerebral volume (Figure 2). The fact that cerebellar tissue loss appears simultaneously with that of the total cerebrum suggests a generalized abnormal process in childhood-onset schizophrenia superimposed on normal development.

When only initial scans were examined, no difference in cerebellar volume was seen between the 50 patients with childhood-onset schizophrenia and the 50 normal comparison subjects. As with other measures in patients with childhood-onset schizophrenia and healthy subjects, only the availability of prospective longitudinal data revealed subtle differences in developmental trajectories (3, 33)

### TABLE 3. Analysis of Developmental Trajectories for Cerebral and Cerebellar Regions in 50 Children With Childhood-Onset Schizophrenia and 50 Healthy Comparison Subjects

<table>
<thead>
<tr>
<th>Structure Measure</th>
<th>Shape of the Curve</th>
<th>Diagnosis Difference in Shape of Curve</th>
<th>Diagnosis Difference in Height of Curve</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>( F ) \text{ df} \text{ } p )</td>
<td>( F ) \text{ df} \text{ } p )</td>
</tr>
<tr>
<td>Total cerebellar volume</td>
<td>Nonlinear</td>
<td>9.30 \text{ 2, 105 } 0.006</td>
<td>1.82 \text{ 1, 97 } 0.18</td>
</tr>
<tr>
<td>Vermal area</td>
<td>No age-related changes</td>
<td>3.63 \text{ 1, 97 } 0.06</td>
<td></td>
</tr>
<tr>
<td>Posterior-inferior volume</td>
<td>No age-related changes</td>
<td>1.10 \text{ 1, 97 } 0.30</td>
<td></td>
</tr>
<tr>
<td>Total cerebral volume</td>
<td>Linear</td>
<td>16.57 \text{ 1, 107 } &lt;0.002</td>
<td>2.69 \text{ 1, 97 } 0.10</td>
</tr>
</tbody>
</table>

*The volumes and areas of male subjects were significantly larger than those of female subjects for all measures: for total cerebellar volume, \( F=48.3, \text{df}=1, 97, p<0.001 \); for total cerebral volume, \( F=55.4, \text{df}=1, 97, p<0.001 \); for posterior-inferior volume, \( F=14.7, \text{df}=1, 97, p<0.001 \); for vermal area, \( F=6.2, \text{df}=1, 97, p<0.02 \). There were no significant sex-by-diagnosis interactions.*

*Derived from mixed model regression.*

**FIGURE 2.** Total Cerebellar and Total Cerebral Volumes in Relation to Age for 108 MRI Scans of 50 Patients With Childhood-Onset Schizophrenia and 101 Scans of 50 Healthy Volunteers

[Graph showing total cerebellar and total cerebral volumes over age for patients and healthy comparison subjects]
that were not detectable in cross-sectional studies with the same subjects (37, 44).

These findings are consistent with reported abnormal cerebellar function in childhood and adult schizophrenia (32, 45–47). There is accumulating evidence for a cognitive role for the cerebellum (see review in reference 48), including executive function and working memory, which are impaired in schizophrenia (49–53). Positron emission tomography studies have revealed abnormalities in cerebellar blood flow while patients carry out cognitive tasks (46, 47). A model of schizophrenia as secondary to disrupted development in a cortico-cerebellar-thalamic-cortical circuit (46) has been termed “cognitive dysmetria” (54, 55), referring to incoordination in the processing, prioritization, retrieval, and expression of information. Unlike a previous study (56), however, we found no significant relationship between any symptom pattern or change in any clinical or cognitive measure and our measures of cerebellar volume loss. We did not have available measures of motor function, which might have revealed a relationship to cerebellar volume loss.

We did not replicate previous findings of decreased midsagittal vermal area and posterior-inferior vermal lobe volume (31), possibly due to the vagaries of hand measurements of small structures. In spite of good intrarater and interrater reliabilities, the anatomy of the vermis is difficult to delineate (40), rater measures are subject to drift, and techniques for hand-measuring are fraught with methodological issues such as location of vermal midslice (39, 57, 58).

Limitations of the study include substantially lower IQ for the patient group. Moreover, since all of our patients were medicated, we cannot rule out the possibility that our findings reflect the effects of neuroleptic medications. Our focus on treatment-resistant subjects also limits the generalizability of these findings.

In summary, the excess loss of brain tissue during adolescence seen in this and other studies may be a trait marker for schizophrenia. Since there is evidence of reduction of cognitive function and working memory, which are impaired in schizophrenia (49–53), we are currently conducting a prospective brain MRI study of the siblings of our patients.

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