Prodromal Symptom Severity Predicts Accelerated Gray Matter Reduction and Third Ventricle Expansion among Clinically High-Risk Youth Developing Psychotic Disorders

Yoonho Chung a  Aron Jacobson a  George He a  Theo G.M. van Erp b  Sarah McEwen c, d  Jean Addington o  Carrie E. Bearden c, d  Kristin Cadenehead e  Barbara Cornblatt f  Daniel H. Mathalon g  Thomas McGlashan h  Diana Perkins k, l  Larry J. Seidman l, j  Ming Tsuang e  Elaine Walker m  Scott W. Woods h  Robert Heinssen n  Tyrone D. Cannon a, h

on behalf of the North American Prodrome Longitudinal Study (NAPLS) Consortium

© 2015 S. Karger AG, Basel

Key Words
Schizophrenia · Psychosis · Prodromal risk syndrome · Magnetic resonance imaging · Scale of Prodromal Symptoms · Clinically high-risk youth

Abstract
A recent prospective longitudinal neuroimaging study of 274 prodromal risk syndrome subjects revealed that those who later developed full-blown psychotic symptoms had exhibited accelerated gray matter loss and third ventricle expansion around the time of psychosis onset. Previous studies also indicate that higher levels of unusual thought content during prodromal states are a significant predictor of psychosis in clinically high-risk (CHR) youth. However, the relationship between clinical symptoms and changes in neuroanatomical structure has not been previously examined in the North American Prodrome Longitudinal Study (NAPLS) sample at the whole-brain level. In this report, we investigated whether symptom severity as measured by the Scale of Prodromal Symptoms (SOPS) predicted the accelerated gray matter decline in 274 CHR cases, including 35 who converted to psychosis. Higher levels of unusual thought content at baseline were associated with a steeper rate of gray matter loss in the prefrontal cortex bilaterally among converters. In contrast, there was no association found among nonconverters. Steeper gray matter loss seems to be unique to those (CHR) individuals with higher levels of subpsychotic predelusional symptoms that acutely worsen in the ramp-up to full-blown psychosis, and as such may reflect pathophysiological processes driving the emergence of psychosis.
**Introduction**

Progressive gray matter reduction in prefrontal regions has been repeatedly observed among clinically high-risk (CHR) youths who develop full-blown psychotic illness [1–8]. As prefrontal cortical regions continue to mature via synaptic pruning and myelination throughout adolescence [9–11], an accelerated rate of synaptic pruning observed among individuals who develop psychosis may explain why the onset of psychosis commonly occurs during late adolescence [12–16]. A recent longitudinal neuroimaging study, with the largest sample of prodromal risk syndrome subjects studied to date (n = 274), revealed that individuals who converted to psychosis showed a steeper rate of gray matter loss in the prefrontal cortex and an increased rate of third ventricle expansion compared with those who did not convert and healthy controls [8]. However, the variability observed in the rate of change in cortical, subcortical, and ventricular structures, even among converters, implicates heterogeneity and individual differences in the pathophysiology of psychosis onset. A question of major importance is whether initial levels of prodromal symptom severity predict a steeper rate of cortical thinning, consistent with the theoretical view that increasing clinical symptom severity during the prodromal state is a consequence of increasing disruption in synaptic activity and functional connectivity in the brain, of which accelerated gray matter loss may be an indicator.

CHR patients are composed of help-seeking individuals with attenuated psychotic symptoms (i.e. psychosis-like symptoms of recent onset that are subpsychotic in intensity, duration, and level of insight) [17]. CHR individuals are considered to be at an imminent risk for psychosis: about 20–25% of them convert to psychosis within 1 year and 30–35% within 2 years [18–20]. Among many established instruments for assessing those at risk, the Scale of Prodromal Symptoms (SOPS) is an operationalized instrument developed and validated to quantitatively assess the severity of prodromal symptoms for psychosis [21–23]. In analyses of psychosis predictors in the first North American Prodrome Longitudinal Study (NAPLS) sample, 2 of the 19 items in the SOPS (i.e. P1 – unusual thought content/unusual ideas; P2 – severity of suspiciousness/persecutory ideas) were most independently predictive of conversion to full psychosis [19]. In particular, higher levels of unusual thought content (such as magical thinking and odd beliefs), rated by P1 symptom criteria in the SOPS, appeared to consistently offer the best prediction among the clusters of symptom ratings in multiple independent samples [19, 24–26].

As a follow-up to the recently published NAPLS longitudinal neuroimaging study [8], we further investigated the relationship between the SOPS P1 symptom severity and neuroanatomical changes in CHR cases in the same NAPLS neuroimaging dataset. We hypothesized that the rate of gray matter reduction in the prefrontal cortex and expansion of the third ventricle would be greater among converters with higher ratings of unusual thought content as measured by SOPS P1 at baseline (BL) assessment.

**Methods**

**Subjects**

The study protocol and consent form were reviewed and approved by the Institutional Review Boards at each of the eight data collection sites (UCLA, Emory University, Beth Israel Deaconess Medical Center, Zucker Hillside Hospital, University of North Carolina, UCSD, University of Calgary, and Yale University). The Structured Interview for Prodromal Syndromes (SIPS) [21] and the Structured Clinical Interview were used for axis I DSM-IV disorders [27] at each assessment. High reliability standards intraclass correlation coefficient (ICC) = 0.92–0.96] for interviewers were met [28]. CHR cases were defined as those who met SIPS/SOPS criteria for a psychosis risk syndrome [21], excluding cases who had ever met DSM-IV criteria for a psychotic disorder. Control participants who met criteria for a psychotic disorder, had a first-degree relative with a current or past psychotic disorder, or met CHR criteria were excluded. Also, general exclusion criteria included substance dependence, neurological disorder, or full-scale IQ <70.

The subjects with MRI structural data included in the analysis are the same cases that were utilized in our previous report, without any additional subjects [8]. Briefly, those with MRI scans were assessed at BL and at the 12-month follow-up (FU). For converters, MRI scans were performed at the point of psychosis conversion. Artifacts in one or both scans led to the exclusion of 14 additional subjects [8]. Briefly, those with MRI scans were assessed at BL and at the 12-month follow-up (FU). For converters, MRI scans were performed at the point of psychosis conversion. Artifacts in one or both scans led to the exclusion of 14 additional subjects [2]. Among 35 CHR cases who converted to psychosis, 239 CHR cases who did not convert, and 135 healthy comparison subjects had usable data and were included. The demographic characteristics of the three groups are shown in table 1.

**MRI Scans**

Five of the data collection sites used Siemens scanners and three sites used GE scanners, all at 3 T. All sites using Siemens scanners employed a 12-channel head coil and all sites using GE scanners employed an 8-channel head coil. Sequence parameters were optimized for each scanner manufacturer, software version, and coil configuration according to the Alzheimer’s Disease Neuroimaging Initiative (ADNI) protocol (http://adni.loni.ucla.edu/research/protocols/mri-protocols/). Scans were acquired in the sagittal plane with a 1 × 1 mm in-plane resolution and a 1.2-mm slice thickness. Siemens scanners used an MPRAGE sequence with a 256 (axial) × 240 (sagittal) × 176 (coronal) mm field of view, TR/TE/TI = 2,300/2.91/900 ms, and a 9-degree flip angle, while GE scanners used an IR-SPGR sequence with a 26-cm field of view, TR/TE/TI = 7.0/minimum full/400 ms, and an 8-degree flip angle.
Image Processing

Identical image processing steps were utilized as in our previous report [8]. All MR images were processed using FreeSurfer version 5.2 (http://surfer.nmr.mgh.harvard.edu/). Surface-based cortical reconstruction was performed to extract thickness measures by calculating the shortest distance from each point on the gray/white boundary to the pial surface at each vertex, along with subcortical volumetric segmentation [29–33]. The subcortical segmentation procedure assigns a neuroanatomical label to each voxel of the MRI volume using a probabilistic atlas and a Bayesian classification rule [31]. The reconstructed BL and FU scans were further processed using FreeSurfer’s longitudinal stream to extract change in thickness and volume estimates [34]. This processing stream initializes each time point scan by utilizing an unbiased within-subject template space and average images [35], created by robust, inverse consistent registration [36], which has been shown to significantly increase statistical power for detecting subtle changes over time [34].

Quality assurance checks were performed to verify all steps in the processing stream. Briefly, this process included visually inspecting the automated reconstruction for skull strip errors, intensity normalization failures, segmentation errors, white and pial surface misplacements, and topological defects. Nearly all scans passed a quality control assessment. A total of 14 subjects, distributed proportionally across the groups, were excluded from the analyses due to unrecoverable MRI artifacts. Control points were utilized to improve surface inaccuracies.

Scale of Prodromal Symptoms

Study participants were evaluated using the SIPS [22] for the presence of one or more CHR syndromes: attenuated subthreshold psychotic symptoms; brief intermittent psychotic symptoms, and substantial functional decline combined with a first-degree relative with a psychotic disorder. The SIPS was administered at initial assessment and at approximately 12 months of FU or at conversion. Based on the SIPS, the severity of symptoms was determined with the SOPS as follows: 0 = absent; 1 = questionably present; 2 = mild; 3 = moderate; 4 = moderately severe; 5 = severe but not psychotic, and 6 = severe and psychotic. The raters achieved high interrater reliability (κ > 0.80), as previously described [19].

Because the ratings ‘1 = questionably present’ and ‘2 = mild’ are considered subprodromal in quality, intensity, and/or frequency, differential gray matter changes among cases with scores of 0, 1, and 2, are not expected. Therefore, we grouped these categories together and rescored them all to 0. Then, syndromal level symptoms such as ‘moderate’ were scored ‘1’, ‘moderately severe’ was scored ‘2’, ‘severe but not psychotic’ was scored ‘3’, and ‘severe and psychotic’ was scored ‘4’ (table 2). This approach yielded a better distribution as there were only 2 converters with ratings in the subprodromal level at BL. Conversion to psychosis was determined by the SIPS criteria that are designed to operationalize the threshold of delusional ideation or hallucination severity required for a DSM-IV psychotic disorder diagnosis [37], as described in detail in prior studies [19, 38].

For the purposes of the present analysis, our focus was on SOPS item P1, indexing unusual thought content, which was the single best predictor of psychosis in NAPLS 1 [19] and also in other in-
Table 2. Detailed description of the rescaled SOPS P1 and P2 symptoms

<table>
<thead>
<tr>
<th>Description</th>
<th>Severity rating</th>
<th>0 = absent/questionably present/mild</th>
<th>1 = moderate</th>
<th>2 = moderately severe</th>
<th>3 = severe but not psychotic</th>
<th>4 = psychotic/extreme</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>P1: Unusual thought content</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perplexity and delusional mood. Mind tricks, such as the sense that something odd is going on or puzzlement and confusion about what is real or imaginary. The familiar feels strange, confusing, ominous, threatening, or has special meaning. Sense that the self, others, or the world has changed. Changes in the perception of time. Déjà vu experience. Nonpersecutory ideas of reference. Overvalued beliefs. Preoccupation with unusually valued ideas. Magical thinking that influences behavior and is inconsistent with subculture norms. Unusual ideas about the body, guilt, nihilism, jealousy, and religion.</td>
<td>0 = no unusual thought content.</td>
<td>1 = probably present/mild</td>
<td>2 = possibly present/mild</td>
<td>3 = definitely present/mild</td>
<td>4 = psychotic or extreme</td>
<td></td>
</tr>
<tr>
<td>P2: Suspiciousness/persecutory ideas</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Persecutory ideas of reference. Suspiciousness or paranoid thinking. Presents a guarded or even openly distrusting attitude that may reflect delusional conviction and intrude of the interview and/or behavior.</td>
<td>0 = no suspiciousness.</td>
<td>1 = probably present/mild</td>
<td>2 = possibly present/mild</td>
<td>3 = definitely present/mild</td>
<td>4 = psychotic or extreme</td>
<td></td>
</tr>
</tbody>
</table>
dependent cohort studies [24–26]. For exploratory purposes, we also examined P2 – severity of suspiciousness/persecutory ideas, which also contributed to psychosis prediction (table 2) [19].

Statistical Analysis

Imaging measures were first transformed to annualized rates of change (ARCH) in each cortical voxel and each subcortical and ventricular region of interest (ROI), where ROIARCH = [(ROIFU – ROIBL)/ROIBL]/interval, and where interval is the time between BL and FU scans in years. Relationships between SOPS P1 and annualized gray matter percent change were tested vertex-wise, bilaterally, across the cortical surface. Monte Carlo simulations, one of the features implemented for multiple comparison correction in FreeSurfer, were used across the surface and synthesized with a cluster-forming threshold of p < 0.05 (two-sided) [39, 40]. To provide a perspective on effects that might be below the statistical threshold, the rescaled P1 ratings were mapped onto the cortical surface with and without multiple comparison corrections.

The annualized rate of change of the third ventricle volume, extracted by FreeSurfer, was chosen as one of the dependent variables, as a significant expansion of this ventricular structure had been observed among converters relative to healthy controls and nonconverters in our previous study [8].

Neither age (p > 0.156) nor sex (p > 0.425) was significantly related to the annualized rate of percent change in the cortical thickness measures and the third ventricle. In addition, age and sex were not significant predictors of change in prefrontal or global cortical thickness within each group [8]. A test-retest reliability study with 8 travelling subjects showed high reliability in overall neuroanatomical measurements [8, 41]. FreeSurfer-derived measures of cortical thickness and subcortical volume in the ROI that we utilized in the secondary analyses showed excellent reliability: superior frontal gyrus (left ICC = 0.908, right ICC = 0.921), rostral middle frontal gyrus (left ICC = 0.842, right ICC = 0.872), and third ventricle (ICC = 0.870). More information on the scanner reliability study can be found elsewhere [8, 41].

Results

Demographic Characteristics

As shown in table 1, there were no significant differences between the diagnostic groups in age, sex, race, or socioeconomic status as measured by income. The converters showed greater BL P1 severity ratings than the nonconverters and healthy controls. In contrast, BL P2 ratings were not significantly different between converters and nonconverters among CHR individuals only. Consistent with previously reported findings [19, 24–26], logistic regression analysis revealed that unusual thought content is a significant predictor of conversion among CHR individuals. For each 1-unit increase in P1 score, the odds ratio of being a converter increases by 2.35 (p < 0.0001). The DSM-IV psychotic disorder diagnosis after the conversion was known for 33 out of the 35 subjects who converted: schizophrenia (n = 8), schizophreniform disorder (n = 9), schizoaffective disorder (n = 2), bipolar disorder with psychotic features (n = 3), brief psychotic disorder (n = 1), and psychosis not otherwise specified (n = 10).

Whole-Brain Analyses

There were no significant differences between converters, nonconverters, and healthy controls in terms of cortical thickness or subcortical or third ventricular volume at BL [8], nor did BL thickness measures vary by SOPS P1 ratings among converters. P1-rescored values at BL were mapped onto the cortical surface maps representing the percent change in cortical thickness (fig. 1).
Clusters of voxels covering the entire rostral middle frontal region (cluster size = 6,828.98 mm², clusterwise p < 0.0001, corrected) in the left hemisphere and the superior frontal region (cluster size = 3,452.78 mm², clusterwise p < 0.0001, corrected) in the right hemisphere survived the whole-brain analysis with cluster-based multiple correction, which showed significant negative correlations with P1 scores at BL. We also display the uncorrected statistical significance maps of the relationship between the annualized rate of change in cortical thickness and SOPS P1 scores, with a nominal p < 0.05 threshold. In the uncorrected maps, in addition to the regions previously identified using cluster-based correction for multiple comparisons, higher P1 ratings at BL predicted steeper rates of gray matter reductions in the right fusiform gyri and in the right paracentral region. The same set of analyses was done on nonconverters and healthy controls; in these groups, no association was found between the clinical symptom ratings and gray matter changes.

For exploratory purposes, we performed the same set of cluster-based whole-brain analyses using the P2 (suspiciousness) severity rating, which was also previously identified as a contributing predictor of psychosis conversion in the NAPLS 1 sample [19]. However, the P2 rating did not show significant associations with the rate of change in gray matter thickness. In addition, we mapped P1 ratings to the rate of change in cortical area and volume using unconstrained vertex maps with a nominal p < 0.05 threshold, but none of the peaks observed survived multiple comparison correction.

**Conversion Status and P1 Interaction**

A multiple regression model estimating an effect of conversion status (converters vs. nonconverters), unusual thought content (P1 rating), and their interaction term on the annualized rate of percent thickness change in the right superior frontal cortex was performed. This model [F(3, 274) = 8.05, p < 0.0001] revealed a strong interaction effect (t = 3.34, p < 0.001), where converters with higher unusual content showed a steeper gray matter decline, whereas nonconverters showed no relationship. The same interaction effect was also present in the left rostral middle frontal cortex [F(3, 274) = 4.21, p < 0.001; P1 × conversion interaction term: t = –3.06, p < 0.01]. The interaction plots for both regions are shown in figure 2.

**Third Ventricle**

The rate of change in third ventricle volume was significantly associated with P1 ratings (r = 0.35, two-tailed p = 0.0289), with higher BL ratings on P1 being associated with a greater rate of third ventricle volume expansion.

---

Chung et al.
**P1 Severity and Duration**

Interestingly, in our previous report, shorter symptom duration was also associated with the steeper gray matter decline and third ventricle expansion among converters [8]. Since both the duration of prodromal symptoms and P1 scores are significant predictors of the steeper gray matter decline and third ventricle expansion among converters, we performed a set of statistical tests that included both the duration of the prodromal state and P1 scores at the BL assessment (within converters) in a set of multiple regression models predicting the rate of change in the regions identified from the whole-brain analysis (i.e. the left rostral middle frontal region and the right superior frontal region). The chosen ROI measures were extracted using the Desikan atlas parcellations [42]. The P1 scores at BL remained nominally significant (p < 0.05 for the right superior frontal gyrus, left rostral middle frontal gyrus, and third ventricle) even after controlling for the duration of prodromal symptoms (table 3), suggesting an independent contribution of the symptom severity in predicting the rate of gray matter decline.

**Discussion**

Higher levels of unusual thought content and a differential rate of prefrontal gray matter reduction have been previously shown to be significant predictors of psychosis in CHR cases [2, 8, 19], but the relationship of these two predictors has not been previously examined in the NAPLS sample at the atlas level. Theoretically, progressive tissue loss in the prefrontal cortex is an indicator of an increasing disruption in integrated synaptic activity and interregional functional connectivity, which are thought to drive the expression of psychotic symptoms. If this is the case, those at risk who show higher levels of positive symptoms at BL should show steeper rates of tissue loss as their symptoms cross the threshold into the psychotic range of severity. Support for this theoretical view was observed in that more severe unusual ideas and beliefs (i.e. mind tricks, external control, magical thinking, etc.) at BL predicted a steeper rate of gray matter loss in the prefrontal cortex bilaterally among the CHR cases progressing to full psychosis. Unusual thought content had previously been shown to be a significant risk predictor of transition to full psychosis in a number of independent datasets [19, 24–26]. According to the SIPS assessment, a prodromal risk syndrome is present when attenuated psychosis-like symptoms have had their onset or worsened in the previous year. Therefore, we can infer that individuals who are rated >1 at the BL assessment are experiencing symptoms that have persisted and intensified since the time of prodromal symptom onset. Since there were no significant differences in cortical thickness, subcortical volume, and ventricular size at BL between converters, nonconverters, and healthy controls [8], and because the neuroanatomical measures at BL did not vary significantly by the initial P1 ratings, the relationships observed in this study suggest a dynamic course of changes in brain structure around the time of psychosis onset.

Interestingly, a significant interaction between the group status and the level of unusual thought content was observed when predicting the rate of change in the prefrontal cortex. When all CHR cases were included in the model, an association between the attenuated positive symptom rating and the neuroanatomical rate of change was observed only among converters. This suggests that an accelerated rate of prefrontal gray matter loss is relevant particularly to CHR individuals whose attenuated positive symptoms acutely worsen over time in association with psychosis onset. It may be that high levels of persistent and intensifying predelusional symptoms are dynamically associated with a reduction in cortical scaffolding that underlies synaptic connectivity and integration of information.

In contrast, the rate of cortical thinning did not differ from that in healthy controls among nonconverters who were characterized by attenuated symptoms that remit over time or remain constant at a subthreshold level. There is likely to be heterogeneity in the causes underlying unusual thought content in the population. It may be that stress or other mechanisms are contributing to the emergence of subpsychotic symptoms among noncon-

<table>
<thead>
<tr>
<th>Predictors</th>
<th>B</th>
<th>SE</th>
<th>t</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Annualized rate of change in the right superior frontal gyrus</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SOPS P1 score</td>
<td>−0.022085</td>
<td>0.008129</td>
<td>−2.149</td>
<td>0.010</td>
</tr>
<tr>
<td>Duration</td>
<td>0.004879</td>
<td>0.002270</td>
<td>2.717</td>
<td>0.039</td>
</tr>
<tr>
<td><strong>Annualized rate of change in the left rostral middle frontal gyrus</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SOPS P1 score</td>
<td>−0.022939</td>
<td>0.010449</td>
<td>−2.195</td>
<td>0.035</td>
</tr>
<tr>
<td>Duration</td>
<td>0.001043</td>
<td>0.002918</td>
<td>0.357</td>
<td>0.723</td>
</tr>
<tr>
<td><strong>Annualized rate of change in the third ventricle</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SOPS P1 score</td>
<td>0.050742</td>
<td>0.023274</td>
<td>2.324</td>
<td>0.026</td>
</tr>
<tr>
<td>Duration</td>
<td>−0.016300</td>
<td>0.006499</td>
<td>−1.615</td>
<td>0.115</td>
</tr>
</tbody>
</table>
stringent p value thresholds were applied with more robust significance on the right when more conversion to psychosis occur in both hemispheres but reported that the differential changes associated with the pronounced in the right hemisphere. Prior studies also progressive tissue loss is occurring bilaterally but is more remained significant after applying a conservative vertex-wise method. This is due to the reduced variation in the clinical ratings assessed at conversion (82% of the cases were rated ±3 on a rescaled P1 rating at conversion). Further, the positive symptom severity rating and the rate of change in gray matter and third ventricle volume were not correlated among nonconverting participants. This is consistent with our previous finding that nonconverters and healthy controls showed no significant group differences in the neuroanatomical developmental trajectories. Taken together, these results indicate that steeper gray matter loss seems to be unique to those individuals with higher levels of persisting and intensifying subpsychotic positive symptoms around the time of psychosis onset. Our findings thus support the development of strategies for the early intervention and care for individuals with high ratings of positive symptoms to either prevent, reduce, or at least delay the disease-promoting processes in the brain.

It is interesting that the correlation between the clinical severity ratings and the cortical changes occurred bilaterally. In our previous report [8], the group differences in the rate of change in cortical thickness in converters versus nonconverters and healthy controls had also shown a bilateral distribution in the prefrontal regions when not correcting for multiple comparisons. However, only the clusters of signals in the right hemisphere had remained significant after applying a conservative vertex-wise correction for multiple comparisons [8, 43]. In this follow-up report on the same sample, we have found that the severity of unusual thought content at BL is associated with bilateral reductions in the prefrontal regions using an anatomically unconstrained vertex-wise method with cluster-based correction for multiple comparisons [39, 40]. This pattern is consistent with the view that the progressive tissue loss is occurring bilaterally but is more pronounced in the right hemisphere. Prior studies also reported that the differential changes associated with the conversion to psychosis occur in both hemispheres but with more robust significance on the right when more stringent p value thresholds were applied [1, 2, 6, 8].

It is worth noting that the steeper rate of gray matter loss in the prefrontal cortex is associated with the P1 symptom ratings (i.e. unusual thought content/unusual ideas) but not with P2 – severity of suspiciousness/persecutory. This may indicate that the emergence of unusual thought content is specifically tied to developmental disturbances in the superior frontal and rostral middle frontal (i.e. dorsolateral prefrontal) regions, and other constellations of psychosis symptomatology may be explained as a secondary phenomenon or hinged on a different pathophysiology. One possible mechanism for explaining the emergence and persistence of delusional beliefs and unusual thought content is based on dysfunctions in belief formation processes, or aberrant prediction error signaling, in which the right prefrontal cortex plays a key role [44, 45]. In addition, it is well known that these prefrontal regions are the neural correlates of top-down regulation and working memory, and functional imaging studies have consistently implicated neural deficits among schizophrenia patients when performing behavioral tasks that require various forms of cognitive control and working memory [46, 47].

The rate of expansion in the third ventricle was significantly associated with BL SOPS P1. Enlarged ventricles have been consistently reported across the literature on schizophrenia [48–50]. The associations of positive symptom severity with the rate of change in both cortical and ventricular structures suggest that the biological underpinnings of the positive symptoms are not limited to the prefrontal cortex, and studies should further examine the disruption of neural circuits at a system level.

Cortical thinning during late adolescence is thought to reflect a pruning and rewiring process via experience-dependent synapse elimination [9]. Hence, a certain degree of thinning in the cortex is expected to be a sign of healthy development. However, it is unclear whether the gray matter decline we observed is an overshoot of a normal pruning process or due to a progressive process specific to schizophrenia per se. Also, based on the design of this study, we cannot conclude whether the progressive tissue loss precedes the onset of full psychosis. As conversions to full psychosis seem to occur in less than 1 year for most of the participants from BL assessments, future studies are encouraged to conduct MRI scans with multiple time points (2- to 3-month intervals) to elucidate whether more precise brain trajectories could predict change in clinical symptoms.

Acknowledgements

The authors thank the following persons for assistance with subject scheduling and/or scan acquisition: Angielette Andaya, Nurit Hirsh, and Jamie Zinberg (UCLA); Richard Juelich (BIDMC...
Disclosed Statement

The authors have no conflicts of interest to declare in relation to the subject of this article. T.D.C. reports that he is a consultant to the Los Angeles County Department of Mental Health on the implementation of early detection and intervention programs for youth at risk for psychosis.

References


