Abnormal cortical folding patterns within Broca’s area in schizophrenia: Evidence from structural MRI

Jonathan J. Wisco a,⁎, Gina Kuperberg a,b,c, Dara Manoach a,b, Brian T. Quinn a, Evelina Busa a, Bruce Fischl a,d,e, Stephan Heckers a,b, A. Gregory Sorensen a,d

a MGH/MIT/HMS Athinoula A. Martinos Center for Biomedical Imaging, Massachusetts General Hospital, Building 149 13th Street, Room 2301, Charlestown, MA 02129, United States
b Department of Psychiatry, Massachusetts General Hospital, Building 149 13th Street, Room 2301, Charlestown, MA 02129, United States
c Department of Psychology, Tufts University, 490 Boston Avenue, Medford, MA 02155, United States
d MIT/HST, Massachusetts Institute of Technology, 77 Massachusetts Avenue, E25-519, Cambridge, MA 02139, United States
e CSAIL, Massachusetts Institute of Technology, The Stata Center, Building 32, 32 Vassar Street, Cambridge, MA 02139, United States

Received 10 October 2006; received in revised form 29 March 2007; accepted 30 March 2007
Available online 9 May 2007

Abstract

We compared cortical folding patterns between patients with schizophrenia and demographically-matched healthy controls in prefrontal and temporal regions of interest. Using the Freesurfer (http://surfer.nmr.mgh.harvard.edu) cortical surface-based reconstruction methodology, we indirectly ascertained cortical displacement and convolution, together, by measuring the degree of metric distortion required to optimally register cortical folding patterns to an average template. An area within the pars triangularis of the left inferior frontal gyrus (Broca’s area) showed significantly reduced metric distortion in the patient group relative to the control group (p = 0.0352). We discuss these findings in relation to the neurodevelopmental hypothesis and language dysfunction in schizophrenia.

© 2007 Elsevier B.V. All rights reserved.

Keywords: Schizophrenia; MRI; Cortical folding; Broca’s area; Language

1. Introduction

During normal cortical development, neurons migrate from the ventricular zone to the cortical plate and begin to establish intrinsic and extrinsic connections around gestational week 20 (Chi et al., 1977; Marin-Padilla, 1992). This occurs with precise timing and is dependent upon a delicate balance of genetic regulation (Rakic, 1995). The connections formed during the prenatal period eventually experience a dramatic period of plasticity in the pre-adolescent years during which pruning occurs (Lewis, 1997). Van Essen hypothesized that, as the brain grows and expands, the strength of
connectivity between neighboring areas of cerebral cortex determines its adult folding patterns such that strongly connected areas form gyri and weakly connected areas form sulci (Van Essen, 1997).

Schizophrenia is a devastating psychiatric illness characterized by positive and negative symptoms as well as severe cognitive impairments across multiple domains including language, memory and executive function. Abnormal cortical development has been thought to be part of the etiology of schizophrenia (Murray and Fearon, 1999; Weinberger, 1987, 1996). Although the pathophysiology of the disease is not yet understood, it has been hypothesized that abnormal structural and/or functional connectivity may lead to its symptoms and cognitive impairments (Elvevag and Weinberger, 2001; Friston, 2005, 1996, 1998, 1999; Friston and Frith, 1995; Weinberger, 1987). Given Van Essen’s hypothesis, it is possible that maldevelopment of intrinsic or extrinsic connections of the cerebral cortex in schizophrenia might manifest as abnormalities of cortical folding patterns in adult patients.

Cortical folding abnormalities in schizophrenia measured using the gyrification index (GI) methodology (Zilles et al., 1988) have been reported in post-mortem (Vogeley et al., 2000) and MRI studies (Harris et al., 1988; Zilles et al., 1988) measured using the gyrification index (GI) methodology. Although the cortex determines its adult folding patterns such that strongly connected areas form gyri and weakly connected areas form sulci (Van Essen, 1997).

Functional connectivity may lead to its symptoms and cognitive impairments (Elvevag and Weinberger, 2001; Friston, 2005, 1996, 1998, 1999; Friston and Frith, 1995; Weinberger, 1987). Given Van Essen’s hypothesis, it is possible that maldevelopment of intrinsic or extrinsic connections of the cerebral cortex in schizophrenia might manifest as abnormalities of cortical folding patterns in adult patients.

Cortical folding abnormalities in schizophrenia measured using the gyrification index (GI) methodology (Zilles et al., 1988) have been reported in post-mortem (Vogeley et al., 2000) and MRI studies (Harris et al., 2004a,b; Kulychny et al., 1997; Sallet et al., 2003). Since the GI measures the ratio of inner and outer cortical surface contours, these studies were limited to analyses of brain slices in different lobes. Cortical folding abnormalities of asymmetry, complexity and variability in the frontal, temporal and parietal lobes of schizophrenia patients have been reported in MRI studies by Narr and colleagues (Narr et al., 2001, 2004). Using cortical surface matching based on major sulcus landmarks, Narr and colleagues found greater variability and deviations of gyral complexity asymmetry in the frontal areas and significant gyral asymmetries in the temporal and parietal regions of schizophrenia patients. All these studies motivated the present study to examine cortical folding within specific, functionally relevant areas of cortex.

We used the Freesurfer cortical surface-based reconstruction methodology (Dale and Sereno, 1993; Dale et al., 1999; Fischl et al., 2001, 1999a,b; Segonne et al., 2004, 2005) to examine cortical folding displacement and convolution, together, in patients with schizophrenia compared to healthy demographically-matched controls. The entire cortical surface of each individual participant was unfolded and the degree of cortical folding was examined within three selected prefrontal and temporal anatomical cortical ROI’s: (1) the left inferior frontal gyrus [Broca’s area, comprised of the pars orbitalis (pOr), pars triangularis (pTr) and pars opercularis (pOp)], and the corresponding regions in the right hemisphere, (2) the right and left dorsolateral prefrontal cortices (DLPFC) and (3) the right and left superior temporal gyri (STG). These cortical regions were selected on the basis of previous morphometric studies that have demonstrated reduced cortical thickness (Kuperberg et al., 2003) and volume (Shenton et al., 2001) in these regions in patients relative to controls, and on the basis of previous fMRI studies reporting that they show abnormal functional modulation and/or functional connectivity in patients relative to controls (Callicott et al., 2000; Jennings et al., 1998; Kuperberg et al., 2006b; Manoach et al., 2000, 1999; Shenton et al., 1992).

The degree of cortical folding was operationalized as the degree of displacement (linear and/or rotational shift) and convolution (complexity of gyrification) of the cortical surface relative to an average template. The template was independently created from the cortical surfaces of a third group of 40 healthy participants (Desikan et al., 2006; Duvernoy, 1991; Ono et al., 1990). Displacement and convolution, together, were indirectly measured from the metric distortion as calculated by the Jacobian energy functional (Fischl et al., 1999a,b) required to align the cortical surface of an individual to the template. We expected cortical surfaces for all individuals to undergo some metric distortion as each unique surface was mapped onto the template, and hypothesized that the degree of metric distortion would be greater in schizophrenia patients than control subjects within each of the a priori ROI’s.

2. Experimental methods

2.1. Participants

Written informed consent was obtained from all subjects before participation according to the established guidelines of the Massachusetts General Hospital Institutional Review Board and the Health Insurance Portability and Accountability Act (HIPAA) guidelines. Twenty-five healthy volunteers (20 male, 5 female) with no history of neurological or psychiatric disorder were recruited by advertisement and 26 patients (22 male, 4 female) diagnosed with schizophrenia according to DSM-IV criteria (American Psychiatric Association, 1990) and on a stable regimen of antipsychotic medication for at least 6 months were recruited from the Erich Lindemann Mental Health Center (Boston, MA). Handedness was measured using the modified Edinburgh Handedness Inventory (Oldfield, 1971; White and Ashton, 1976). Patients and controls were gender, age and handedness matched (see Table 1).
Exclusion criteria included neurological disease or damage, head trauma with documented cognitive impairment or loss of consciousness greater than 5 min, medical disorders that could impair neurocognitive function, a history of substance abuse within three months prior to participation and specific exclusions for MRI.

2.2. MRI acquisition

We acquired two data sets of T1-weighted MPRAGE images as part of a larger set of structural data that were acquired during a 1.5 h scan session. The MPRAGE protocol was run in a Siemens 3T Trio scanner (Siemens Medical Solutions, Erlangen, Germany) using an 8-channel phased-array head coil: TR/TE/TI=2530/3.45/1100 ms; flip angle=7°; FOV =256 mm; 1.3×1.0×1.3 voxels; 128 slices in the sagittal plane. Images covered the entire brain. Motion artifact was monitored throughout the scan session. The protocol for any image data set that exhibited excessive motion was re-run after verbally informing the participants.

2.3. Image reconstruction

Individual cortical surfaces were reconstructed using the Freesurfer analyses tools (http://surfer.nmr.mgh.harvard.edu) developed at the Martinos Center for Biomedical Imaging as described in detail previously (Dale and Sereno, 1993; Dale et al., 1999; Fischl et al., 2001, 1999a,b; Segonne et al., 2004, 2005). First, image intensity variations due to magnetic field inhomogeneities were normalized. Second, extra-cerebral voxels were removed, essentially "skull-stripping" the brain. Third, the resulting intensity normalized, skull-stripped brain images were segmented to classify voxels into gray and white matter regions. Fourth, cutting planes were computed to separate the cerebral hemispheres from each other and from the brainstem and cerebellum. Finally, the resulting hemisphere volumes were covered with a triangular tessellation and deformed to produce a representation of the gray/white matter interface for each subject. Any geometric inaccuracies were corrected manually and the reconstruction procedure was repeated until an accurate surface was achieved. The gray/white matter interface surface was inflated and transformed into a spherical representation (Fig. 1).

2.4. Measuring metric distortion

Each individual subject’s spheres were non-linearly registered to an average template sphere created from 40 independent control subjects (see example in
Fig. 1. Cortical reconstruction. A given subject brain represented here by a 3D pial surface reconstruction (a) was transformed into gray matter/white matter interface surface (b), inflated (c) and transformed again into a spherical surface (d). Gyri and sulci are represented in light gray and dark gray colors, respectively.

Fig. 2) (Fischl et al., 1999a). A spherical coordinate system based on the average template was translated back to the subject spheres such that every coordinate corresponded across all subjects, and thus every participant’s brain was registered into a common coordinate system.

In order to achieve the registration of individual spheres into the common coordinate system, the triangular
tessellation used to represent the gray/white matter interface and its spherical representation was necessarily deformed. The deformation of a particular tessellation vertex was its metric distortion, which was calculated using the following general equation:

\[
\text{Metric distortion} = \frac{k \times \text{area of a triangle on registered sphere}}{\text{area of triangle on original gray/white interface surface}}
\]

Where \( k \) is the ratio of the total surface area of the original gray/white interface surface to the total surface area of the individual sphere.

We measured the metric distortion required to achieve optimal spherical registration of gyri and sulci (at each tessellation vertex) across individual cortical surfaces (Fischl et al., 1999b). We restricted statistical analysis within three ROI’s that were generated on each subject’s gray/white matter interface surface during the Freesurfer reconstruction stream (see Fig. 3). [These ROI’s were validated previously (Desikan et al., 2006; Duvernoy, 1991; Ono et al., 1990).] The ROI’s included: (1) The inferior frontal gyri [pars orbitalis (pOr), pars triangularis (pTr) and pars opercularis (pOp) gyri, comprising Broca’s area on the left] operationally defined as the three adjacent gyri rostral to the precentral gyrus (Desikan et al., 2006; Duvernoy, 1991; Ono et al., 1990); (2) bilateral dorsolateral prefrontal cortices (DLPFC), operationally defined as the middle third of the middle frontal gyrus (Rajkowska and Goldman-Rakic, 1995); (3) bilateral superior temporal gyr (STG) operationally defined as the gyrus surrounded by the Sylvian fissure, superior temporal fissure and supramarginal gyrus (Desikan et al., 2006; Duvernoy, 1991; Ono et al., 1990).

2.5. Statistical analysis of metric distortion

We used a General Linear Model (GLM) to calculate cohort differences in metric distortion at each tessellation vertex within the a priori ROI’s (inferior frontal gyrus, DLPFC, STG) of both hemispheres. Monte Carlo permutation cluster analyses were then performed to correct for multiple comparisons using a cluster threshold of 0.0500. Finally, the mean metric distortion was calculated in any clusters that survived multiple comparisons analysis.

3. Results

In the left hemisphere, clusters that were significant \((p<0.05)\) before correction for multiple comparisons were located in pars triangularis (pTr) (maxima \(p_1=0.0030\) and pars opercularis (pOp) (maxima \(p_1=0.0039, p_2=0.0481\)) (Table 2). No clusters were detected in the pars orbitalis (pOr), the left dorsolateral prefrontal cortex (L-DLPFC) or left superior temporal gyrus (L-STG). The only cluster surviving correction for

![Fig. 3. Cortical folding variability in Broca’s area (pOr, pTr, pOp), dorsolateral prefrontal cortex (DLPFC) and superior temporal gyrus (STG). This illustrates the individual cortical folding variability of these ROI’s between two control subjects (a, b) and two schizophrenia patients (c, d) in a pial surface reconstruction of brain image volumes. In the spherical registration of these subjects to an average subject template, the metric distortion required for optimal alignment is a reflection individual cortical displacement and convolution.](image-url)
multiple comparisons was located in pTr (630.18 mm², clusterwise p = 0.0352). Metric distortion in this cluster was greater in schizophrenia patients relative to control subjects (Fig. 4). Within the pTr cluster, we found the mean (SD) metric distortion in schizophrenia patients was 1.579 (0.269) and in control subjects was 1.377 (0.225).

In the right hemisphere, clusters that were significant before multiple comparisons correction were located in pOr (maxima p = 0.0071), pTr (maxima p = 0.0041) and pOp (maxima p = 0.0042) (Table 2). No clusters were found in the R-DLPFC or the R-STG. None of the clusters in the right hemisphere survived correction for multiple comparisons.

4. Discussion

4.1. Cortical development and schizophrenia

Previous studies have shown regional cortical folding abnormalities of gyrification (Harris et al., 2004a,b; Kulynych et al., 1997; Sallet et al., 2003; Vogeley et al.,...
asymmetry, complexity and variability (Narr et al., 2001, 2004) in patients with schizophrenia. These studies used a variety of methods to compare the spatial distribution of entire gyri and sulci. Their results motivated the present study to measure metric distortion at each point on the tessellated surface of the left inferior frontal gyrus, dorsolateral prefrontal cortex (DLPFC) and superior temporal gyrus, thereby providing an analysis of folding abnormalities within these cortical areas.

We have thus demonstrated an area within the left pars triangularis (pTr) of Broca’s area in which metric distortion was significantly greater in schizophrenia patients relative to healthy, demographically-matched control participants. In the spherical registration algorithm of the Freesurfer analysis tools, the energy functional to generate accurate alignment includes a term for minimizing metric distortion while optimally aligning sulci and gyri (Fischl et al., 1999a,b). The magnitude of metric distortion, therefore, is an indirect measurement of cortical displacement and convolution, together, relative to the average subject template.

The cerebral cortex undergoes marked changes in its intrinsic and extrinsic connectivity over the life span (Chi et al., 1977; Lewis, 1997; Marin-Padilla, 1992; Rakic, 1995). Broca’s area, specifically, is known to undergo substantial plasticity from birth to adulthood (Amunts et al., 2003; Etchepareborda and Lopez-Lazaro, 2005). The genetic process controlling cortical development may contribute to the onset and expression of schizophrenia (Woolf, 1997; Wyatt, 1996).

Volume measurement has been used as another index for assessing the integrity of gray matter and white matter. Bartzokis and colleagues made an interesting observation of white matter volume increase with concomitant gray matter volume decrease between the second and fourth decades of life in the frontal lobes of normal aging subjects (Bartzokis et al., 2001). However, within the same age range, they found an absence of white matter volume expansion accompanying gray matter volume reduction in the frontal lobes of patients with schizophrenia (Bartzokis et al., 2003). These studies together suggested that normal white matter myelination with increasing age is abnormal in patients with schizophrenia. White matter atrophy in the frontal lobes has been noted in other recent studies (Sanfilipo et al., 2000; Wible et al., 2001), but the discrepancy may be due to not taking the normal aging process into account.

Volume changes due to normal aging and the disease process of schizophrenia may affect the relative geometric surface patterns of sulci and gyri. Although Kuperberg and colleagues showed cortical thinning in specific areas of the frontal lobe, including Broca’s area (Kuperberg et al., 2003), these changes in gray matter volume are unlikely to alter the relative position of sulci or gyri, which are more likely determined by the geometry of the underlying white matter (Van Essen, 1997). Aberrant white matter development between the second and fourth decades of life in schizophrenia, as observed by Bartzokis et al. (2003), would contribute most to abnormal cortical folding patterns. After the fourth decade, however, degeneration of white matter due to normal aging would contribute more to changes in cortical geometry. In this study, we measured metric distortion during the fourth decade of life (mean age of subjects and patients was approximately 45 years, see Table 1), which would be between the time gray matter and white matter development would be ending and normal aging degenerative processes would be beginning. Therefore, the results presented here may reflect the disease process of schizophrenia before degeneration of normal aging.

4.2. Functional significance of abnormalities in Broca’s area

The precise functional significance of cortical folding abnormalities in Broca’s area in schizophrenia is unclear. However, given that abnormal cortical folding may reflect abnormal structural and/or functional cortical connectivity, we can speculate on potential functional links between cortical folding abnormalities in this area and the symptoms and patterns of cognitive dysfunction that characterize the schizophrenia syndrome.

In the healthy brain, the left inferior prefrontal gyrus is a major component of the circuitry mediating both language comprehension and language output. Its precise role is debated but it has been implicated in aspects of semantic processing including semantic encoding (Demb et al., 1995; Kirchhoff et al., 2000; Wagner et al., 1998), semantic priming (Copland et al., 2003; Giesbrecht et al., 2004; Kotz et al., 2002; Kuperberg et al., 2006b; Matsumoto et al., 2005; Wheatley et al., 2005), controlled semantic retrieval (Wagner et al., 2001), and semantic selection (Fletcher et al., 2000; Moss et al., 2005; Thompson-Schill et al., 1997, 1999). Broca’s area has also been implicated in phonological processing (Poldrack et al., 1999), aspects of syntactic processing (Caplan, 2001; Caplan et al., 2000), syntactic working memory (Fiebach et al., 2005) and, more generally, in other aspects of verbal working memory function (Barde and Thompson-Schill, 2002), and in speech output (Blank et al., 2002).

Schizophrenia has long been characterized as a disorder of thought, language and communication (Andreasen, 1979; Bleuler, 1911/1950). Many of its core symptoms including verbal hallucinations and positive thought
disorder with disorganized speech are characterized by language dysfunction (American Psychiatric Association, 1990). Cognitively, language deficits have been identified in children at risk for schizophrenia (Cannon et al., 2002; Fuller et al., 2002; Ott et al., 2001) as well as in patients in their first episode of psychosis (Fuller et al., 2002; Hoff et al., 1999). Numerous neuropsychological and psycholinguistic studies have established that patients with schizophrenia show selective language abnormalities as they monitor the source of verbal input (Ditman and Kuperberg, 2005; Johns et al., 2001), explicitly or implicitly retrieve information from semantic memory (Kuperberg, 2005; Johns et al., 2001), explicitly or implicitly retrieve information from semantic memory (Kuperberg et al., 2007), although abnormal function in this region is not usually seen in isolation, but rather in association with abnormal modulation of other language regions including superior, inferior and medial temporal cortices.

It is possible that localized abnormalities in folding within Broca’s area reflect abnormal cortical connectivity in this region and that this, in turn, leads to widespread cortical dysfunction during language processing in schizophrenia and to the clinical language disturbances that characterize this disorder. However, the current sample size was relatively small and, at uncorrected thresholds, other regions including the inferior parietal lobule and frontopolar region (not reported here) also showed abnormal folding. Future studies with larger samples will determine the degree to which cortical folding abnormalities are localized or generalized across the cortical ribbon, and whether these abnormalities predict the severity of specific symptoms or specific cognitive deficits in schizophrenia.

4.3. Caveats

The present study included 26 patients and 25 demographically-matched healthy controls, a sample size that would be adequate for statistical analysis of gross morphological changes. However, we measured cohort differences in metric distortion at several thousand tessellation vertices, even within a priori ROI’s. Table 2 illustrates the stringent statistical analysis performed to avoid false-positive results. The most plausible solution leading to the discovery of more clusters of significant metric distortion that might be expected in other areas of the cortex responsible for language would be to add many more subjects.

The sample of patients for this study was limited to chronic, medicated patients. The present study cannot account for whether medication might also contribute to the metric distortions associated with the disease. An opportunity to measure metric distortion in non-medicated patients would be valuable to pursue.

Acknowledgements

Support for this research was provided in part by the Athinoulia A. Martinsos Foundation, GlaxoSmithKline, the National Cancer Institute (ST32 CA09502), the National Institute for Mental Health (RO1 MH071635, R01 MH067720), the National Center for Research Resources (P41-RR14075, R01 RR16594-01A1 and the NCRR BIRN Morphometric Project BIRN002, U24 RR021382), the National Institute for Biomedical Imaging and Bioengineering (R01 EB001550), the National Institute for Neurological Disorders and Stroke (R01 NS052585-01) as well as the Mental Illness and Neuroscience Discovery (MIND) Institute. Dr. Fischl is also part of the National Alliance for Medical Image Computing (NAMIC), which is funded by the National Institutes of Health through the NIH Roadmap for Medical Research, Grant (U54 EB005149). Information on the National Centers for Biomedical Computing can be obtained from http://nihroadmap.nih.gov/bioinformatics.

The authors would like to thank Mary Foley, Michael Zussman Christine Portal and Stuart Wallace for their invaluable assistance with the imaging sessions, recruitment and management of patients. We would like to thank Jenni Pacheco for her helpful advice with the cortical surface reconstructions. Finally, we would like to thank Dr. Doug Greve for his statistical advice.

References


Weinberger, D.R., 1996. On the plausibility of “the neurodevelopmental hypothesis” of schizophrenia. Neuropsychopharmacology 14, 1S–11S.


