The subgenual cingulate gyrus exhibits lower rates of bifurcation in schizophrenia than in controls, bipolar disorder and depression

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ABSTRACT

The subgenual cingulate cortex has been found to be different in structure and function in mood and affective disorders compared to healthy individuals. Imaging studies have shown a decrease in function of the subgenual region in bipolar disorder and depression, with overall glial number shown to be decreased in these disorders. Decreases in subgenual grey matter in SZ have been observed also. In this neuropathological study upon formalin-fixed coronal brain sections we describe the morphological finding of decreased frequency of subgenual cingulate crown bifurcation (p = 0.02) as compared to control, bipolar and depression cases. This suggests that the cingulate cortex in schizophrenia may be morphologically distinct in utero formation, potentially enabling an early identification of high-risk individuals.

Keywords: Neuropathology; Schizophrenia; Cortical; Flattening

1. INTRODUCTION

The subgenual cingulate cortex (SCC), part of Brodmann area 24a, is a region of interest in schizophrenia (SZ) and bipolar disorder (BPD). The anterior cingulate cortex (ACC) lies on the medial surface of the cerebral hemisphere, covering the anterior part of the corpus callosum. It is defined as Broadmann area 24a. The ACC extends from the subgenual ventral terminus and continues rostral to the genu of the corpus callosum following the dorsal surface.

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issue with the morphology of anterior cingulate in these disorders.

In this study we have taken coronal sections through the SCC in cases of SZ, BPD and RDD and control cases to examine the prevalence of cortical bifurcation. The size of the bifurcation within a gyrus means it is not visible by imaging and therefore requires a microscopic methodology. As the extent of folding of the cortex increases its surface area within a given gyral volume, potentially reflecting the number of cortical neurons, this pattern of morphology may reflect a smaller cortical volume present from initial neurodevelopment of the cingulate.

2. METHODS

A cohort of cases obtained from the Corsellis brain collection was used in neuropathological investigation of the cingulate in a series of studies (Williams, 2012). Details of the diagnostic groups are shown in Table 1. A total of 68 age and post-mortem interval (PMI) matched cases were used, with 20 control cases with no psychiatric disorder (NPD), 12 SZ cases, 16 BPD cases and 20 cases of RDD.

All cases underwent full neuropsychiatric review and were subject to full neuropathological screening. Cases involving heavy alcohol or drug use and those exhibiting significant pathology such as neurodegenerative disease, cerebral vascular disease, ischemic brain damage, CNS infections and traumatic brain injury were excluded from the study. The majority of samples come from the county of Essex but a smaller number came from national referrals [27]. Medical notes were reviewed by a consultant psychiatrist and patients were selected on fulfilment of the ICD-10 criteria for SZ and MDD, for robustness of diagnosis the brains of the three diagnostic groups all suffered from chronic illness. Assessment for any neurodegenerative, neurovascular or infectious pathology, including Parkinson’s disease, was undertaken by a consultant neuropathologist and affected patients excluded. Patients with any recorded history of alcohol or drug abuse were also excluded. In SZ selection the presence of first-rank symptoms was a necessity, and cases with onset younger than 20 and older than 30 were excluded.

Bilateral SCC and adjacent corpus callosum were dissected from formalin-fixed coronal blocks by a consultant neuropathologist and immersed in 10% formalin (4% formaldehyde v/v) until processing. Processing involved serial immersion in formalin, alcohols, methanol and xylene overnight. Tissue blocks were then embedded in paraffin wax and stored at 4°C. Paraffin-embedded blocks were serially sectioned in the coronal plane at 10 µm and mounted on 25 × 75 mm electrostatic glass slides. Slides were blinded by an investigator not involved with the project before measurement. Sections from within 100 µm each case were stained with hematoxylin and eosin (H & E) and cresyl-violet (CV) for accurate examination of the anatomy of the structures. Sections for H & E stain were submerged in xylene for 30 minutes and serially placed in troughs of 2 × 100%, 90%, 70% ethanol (EtOH) and pure distilled water, each for 2 minutes. Slides were immersed in Mayer’s haematoxylin stain for 5 minutes and subsequently washed in distilled water. Specimens were placed in differentiating agent acid-alcohol (1% HCl/70% EtOH) for 10 seconds and into distilled water for 2 minutes. Sections were then immersed in 1% eosin stain for 5 minutes and washed in distilled water. Finally, specimens were dehydrated in serial baths of 70%, 90% and 2 × 100% IMS for 2 minutes each and submerged in xylene for 20 minutes before cover slips were fixed using DPX. Sections for CV stain were immersed in xylene for 30 minutes, incubated in 100%, 90% and then 70% alcohol for 10 minutes each, before immersion in ultra-pure water. Sections were immersed in cresyl-violet stain for 5 minutes before washing in ultra-pure water and differentiation in 95% alcohol/ethanoic acid. After washing in ultra-pure water sections were dehydrated in serial alcohols, immersed in xylene and mounted with DPX.

Images of both H & E and CV stained coronal sections were taken using an Olympus microscope at 40× total magnification and captured at 2096 × 1536 resolution covering 2200 × 1600 µm area. Image analysis was performed using Image Pro Plus software (Media Cybernetics, US), calibrated using an optical graticule. In total for regions were measured in each case, both left and right SCC in H & E and CV sections. H & E sections were consulted to define the limits of the SCC grey matter. Measures were taken in CV and H & E stained sections

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>n</th>
<th>Age/y</th>
<th>Sex ratio M/F</th>
<th>PM Delay/h</th>
<th>Fixation time/y</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPD</td>
<td>20</td>
<td>65.5(2.34)</td>
<td>12/8</td>
<td>44.2(7.38)</td>
<td>10.2(0.52)</td>
</tr>
<tr>
<td>SZ</td>
<td>12</td>
<td>58(6.44)</td>
<td>6/6</td>
<td>47.8(10.5)</td>
<td>11.1(1.30)</td>
</tr>
<tr>
<td>BPD</td>
<td>16</td>
<td>56.1(5.21)</td>
<td>7/9</td>
<td>50.5(7.46)</td>
<td>19.4(2.02)</td>
</tr>
<tr>
<td>RDD</td>
<td>20</td>
<td>47.6(3.12)</td>
<td>6/14</td>
<td>37.3(5.62)</td>
<td>10.4(0.88)</td>
</tr>
</tbody>
</table>

Table 1. Summary group data. NPD—No Psychiatric Disorder; SZ—Schizophrenia; BPD—Bipolar Disorder; RDD—Recurrent Depressive Disorder. Age, PM delay and fixation time shown as means with SEM in brackets.
by a scoring of incidence of bifurcation in the SCC from both hemispheres from each case, after comparable measures were taken from adjacent sulci and from the level of the crown. Bifurcation was defined as the existence of a sulcal-type projection into the crown of the primary SCC that were less than 50% the depth of both the callosal-cingulate sulcus and the primary cingulate sulcus, shown in Figure 1. Scoring of bifurcation of was taken from both left and right primary SCC in both CV and H & E slides, totalling two measures per hemisphere and four measures per case. Bifurcation had to be present in both CV and H & E slides for inclusion.

Measures were performed blind to diagnosis and were unblinded by an investigator not involved with the project before analysis using Image Pro Plus. Multiple comparisons of confounding variables were performed using the general linear model univariate analysis using SPSS v16.0 statistical software (SPSS, USA). Direct analysis of the incidence of bifurcation was performed using Pearson's Chi-Square test (SPSS, USA).

3. RESULTS

The MDD group had a significantly lower age of death as compared to controls (p = 0.038, 1-way ANOVA). The MDD group (n = 20) contained 11 confirmed cases of suicide. The other disease groups also showed instances of suicide, which may contribute toward the non-significant trend downward in age in SZ (1 suicide, n = 12) and BPD (3 suicides, n = 16). The brain tissue of the BPD group was in formalin significantly longer than the other groups (Mean fixation time: NPD 11.3 yr vs. BPD 19.7 yr, p = 0.0004; NPD vs. SZ. 11.6 yr, p = 0.074; NPD vs. RDD. 11.8 yr, p = 0.59, ANOVA). There was no effect of sex, age, PM delay, incidence of suicide, fixation time or hemisphere on the incidence of bifurcation.

The incidence of bifurcation was 11/20 NPD cases (55%), 2/12 SZ cases (17%), 9/16 BPD cases (56%) and 11/20 RDD cases (55%), shown in Figure 2. The SZ group showed a significantly lower incidence of bifurcation than the controls or other diagnostic groups (p = 0.02, Pearson’s Chi-Square test, SPSS v16.0).

4. DISCUSSION

The results suggest that SCC bifurcation is less common in schizophrenia than in either controls, BPD or RDD. Whether this bifurcation is related to changed numbers of neuron or glial cells, or the function of the SCC, are unknown. Future neuropathological studies will be required to elicit further information of the cellular changes associated with altered cortical folding. Due to the nature of neuropathological studies it is not possible to measure total SCC volume using this type of study.

The cases of bipolar disorder had a longer mean PM delay than the cases from the other diagnostic groups. This was predominantly due to three male bipolar cases, numbers 32 (89 h), 34 (91 h) and 45 (100 h). The RDD group showed a reduced age of death, which may reflect a higher number of suicides amongst this group. Similarly male schizophrenia cases showed a younger age of death than female schizophrenia cases which was likely related to a higher number of suicides. Additionally the BPD cohort has a longer period of fixation, due to the lower number of BPD cases in the tissue bank, requiring tissue from earlier donations to be included. However these variables had no effect on the occurrence of bifurcation, or previous variables measured in the cingulate of these cases [21,28]. We did not have data on the illness duration. However as a requirement for inclusion was symptom onset between 20 - 30 yr this strongly correlated age, which had no effect on incidence of bifurcation.

Although the measures were collected from slides within 100 μm of one another bifurcation was a gross
anatomical occurrence and was easily visible with the naked eye on the blocks. Ideally the full volume of the SCC could be estimated by many serial sections, but this would require many blocks along the entire structure and this amount of tissue was not available. Also there would be errors involved due to the tissue shrinkage during fixation and processing. If these technical issues could be overcome then a volumetric study would be extremely useful as the measurements could be performed with far greater accuracy than in imaging.

Cortical folding changes in length and depth of temporal and frontal lobes have been reported in schizophrenia [29]. These changes been shown to be present before disease onset. Patients with adolescent onset schizophrenia have significantly more flattened curvature in the sulci and more steeped or peaked curvature in the gyri [30], and increased cortical folding in the superior frontal cortex and in gyral and sucal folding in temporal lobe of first episode schizophrenia [31,32]. High risk individuals have been observed to have altered cortical folding in the prefrontal cortex [33]. Examination of bifurcation may give additional information of changes in the cortex in schizophrenia between the scales of neuropathological reports of cell density and morphometry and the imaging data showing larger scale trends across cortical regions.

As gyral morphology is created in utero during the initial folding of the cortex, this suggests that morphological changes in schizophrenia may be hardwired early in life. If these can be identified then they may help with the early identification of high-risk individuals. Whilst large-scale changes in gyral folding may only be detectable in detailed analysis of the whole brain surface this study suggests that by looking at regions of high vulnerability signs of this change may be measurable at the microscopic level.

5. ACKNOWLEDGEMENTS

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REFERENCES


