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Stereological assessment of the dorsal anterior cingulate cortex in schizophrenia: absence of changes in neuronal and glial densities

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Abstract

Aims—The prefrontal and anterior cingulate cortices are implicated in schizophrenia, and many studies have assessed volume, cortical thickness, and neuronal densities or numbers in these regions. Available data however are rather conflicting and no clear cortical alteration pattern has been established. Changes in oligodendrocytes and white matter have been observed in schizophrenia, introducing a hypothesis about a myelin deficit as a key event in disease development.

Methods—We investigated the dorsal anterior cingulate cortex (dACC) in 13 males with schizophrenia and 13 age- and gender-matched controls. We assessed stereologically the dACC volume, neuronal and glial densities, total neuron and glial numbers, and glia/neuron (GNI) ratios in both layers II-III and V-VI.

Results—We observed no differences in neuronal or glial densities. No changes were observed in dACC cortical volume, total neuron numbers, and total glial numbers in schizophrenia. This contrasts with previous findings and suggests that the dACC may not undergo as severe changes in schizophrenia as is generally believed. However, we observed higher glial densities in layers V-VI than in layers II-III in both controls and patients with schizophrenia, pointing to possible layerspecific effects on oligodendrocyte distribution during development.

Conclusions—Using rigorous stereological methods, we demonstrate a seemingly normal cortical organization in an important neocortical area for schizophrenia, emphasizing the importance of such morphometric approaches in quantitative neuropathology. We discuss the significance of subregion- and layer-specific alterations in the development of schizophrenia, and the discrepancies between post-mortem histopathological studies and *in vivo* brain imaging findings in patients.

Keywords

dysmyelination; oligodendrocytes; white matter; morphology; cytoarchitecture; myelin

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INTRODUCTION

White matter deficits and oligodendrocyte pathology have gained increasing interest in schizophrenia research, in light of findings that showed changes in expression levels of myelin and oligodendrocyte-related genes [1–3], and previous imaging findings showing alterations in the white matter [4, 12]. The myelin hypothesis in schizophrenia has been extensively reviewed in [13–18]. In the present study, we further investigate the role of a myelin deficit, in the context of possible neuropathology of the anterior cingulate cortex (ACC) in schizophrenia.

The ACC has been much studied in schizophrenia. It is known to play a role in the regulation of cognition, attention, emotions, and in social interactions. In the human brain, the ACC can be subdivided along its rostrocaudal and dorsoventral axes, through which there exist gradients in cytoarchitecture and topography in its afferent and efferent projections [19–22]. The ACC consists of areas 24 and 25 surrounding the genu of the corpus callosum. Some authors also include the paracingulate area 32 in the ACC. Cytoarchitecturally the ACC is complex and can be divided into distinct subregions [19, 21– 23], and displays large individual variability and hemispheric asymmetry in the human brain [19]. The ACC receives multimodal sensory information from insular, temporal, and parietal association cortices, and emotional information from the amygdala and the orbitofrontal cortex [24–28]. It shows extensive connectivity with the dorsolateral prefrontal cortex. Specifically, the dorsal ACC (dACC, or supragenual part) is connected with the dorsolateral prefrontal and premotor areas, while the rostral ACC (rACC, or subgenual and pregenual parts) is connected with the amygdala and the orbitofrontal cortex [29]. These different connectivity patterns establish two functional ACC domains, a dorsal cognitive subdivision and a ventral affective subdivision, as demonstrated by imaging studies [30–32]. Based on its connections with other areas, the dACC (corresponding to subdivisions 24a'b' and c') is involved in attention to action, error detection, and motor functions [30], while the rostral ACC (corresponding to subdivisions 24abc, area 25 and the paracingulate area 32) is involved in the evaluation and regulation of emotion [33, 34].

Stereologicalal methods have been used to investigate the oligodendrocyte densities and distribution patterns in the dorsolateral prefrontal and anterior cingulate regions. Oligodendrocyte densities have previously been found to be decreased in schizophrenia in white and grey matter in area 9 [35], and in area 24 [36], but not in area 32 [36], or in the cingulum bundle [37]. Morphometric studies of neuron densities have been made in a limited number of cortical regions, and investigators have observed decreases in neuronal densities in area 24 [38], while increases have been found in areas 9 and 46 [39, 40]. However, using unbiased stereologicalal approaches, these findings have not been replicated. Moreover, brain imaging findings frequently reporting decreases in cortical or white matter volumes [41–45] are difficult to correlate to volume estimates from neurohistological methods. To investigate further the neuropathology of neurons and glia in schizophrenia in the ACC region, we studied the dACC using stereologicalal approaches.

MATERIALS AND METHODS

Subjects

This study was performed on the same post-mortem brains (both hemispheres) from 13 male patients with schizophrenia and from 13 age-matched male controls that were investigated in our previous studies on neuron numbers in subcortical regions [46], capillary length densities in the frontal cortex [47], and mean neuron spacing abnormalities in the neocortex [48].

All cases with schizophrenia had been patients either in German university hospitals or in German State psychiatric hospitals and full clinical records were available. All control cases had been patients either in German university hospitals or in German local district hospitals. Patients with schizophrenia and controls were similar in terms of ethnic background. In all of the cases, autopsy was performed after consent was obtained from a relative according to the laws of the Federal Republic of Germany. The use of autopsy cases for scientific investigations was approved by the relevant institutional review boards and ethics committees. Records from autopsy (including a summary of the medical history) were available for all patients with schizophrenia, and for all controls. All patients with schizophrenia met the diagnostic criteria of the Diagnostic Statistical Manual, DSM-IV [49] and International Statistical Classification of Diseases and Related Health Problems, ICD-10 [50]. All patients with schizophrenia were subjected to long-term treatment with typical neuroleptics. However, due to the fact that most of the patients were not hospitalized throughout the duration of their illness, the clinical records did not cover fully the entire medication histories and it was therefore not possible to calculate lifetime medication exposures. The clinical notes were assessed by two experienced clinical psychiatrists for clear evidence that the diagnosis of patients with schizophrenia was concordant with DSM-IV criteria for schizophrenia, and to ensure that the brains from the controls were free from psychopathology.

All pathologists involved in the autopsies were instructed by H.H. and adhered to identical handling and processing conditions of the brains. Brains were fixed by immersion in 4% formalin (one part 40% aqueous formaldehyde in nine parts H_2O) ranging from three months to ten years prior to histological processing as previously described [51]. The age of the patients, clinical diagnoses, illness duration, causes of death, the post-mortem interval and the fixation time are summarized in Table 1. The patients with schizophrenia did not differ from the controls with respect to mean age (two-tailed Student's t-test; $P = 0.946$), mean post-mortem interval ($P = 0.581$) and mean fixation time ($P = 0.089$). The mean age at onset of the patients with schizophrenia was 22.9 ± 1.5 years.

Exclusion criteria for both patients with schizophrenia and the controls comprised neurological problems that required intervention or interfered with cognitive assessment (e.g., stroke with aphasia), history of recurrent seizure disorder, history of severe head injury with loss of consciousness, history of other psychiatric illness, history of self-administered intoxication, and diabetes mellitus with free plasma glucose >200 mg/dl. After histological processing of the brains, each section was coded and verified for the absence of tumours, infarcts, heterotopias, signs of autolysis, staining artefacts and gliosis. In addition, from all the brains of the schizophrenic patients and controls older than 40 years, sections through the central portion of the entorhinal and transentorhinal cortex that were not stained with gallocyanin (see details below) were labelled with the Gallyas method to detect neurofibrillary changes [52]. Neurofibrillary tangles were very rarely detected in the transentorhinal and entorhinal cortex on Gallyas-stained sections, compatible with Braak's stage I [53].

Tissue processing

The brainstem with the cerebellum was separated from the forebrain at the level of the rostral pons, and the hemispheres were divided medio-sagittally. Both hemispheres were then cut into serial 600–700µm-thick coronal sections as previously described [51]. Briefly, the hemispheres were cryoprotected in a mixture of 2% glycerol-dimethylsulfoxide/ 4% formaldehyde/ 10–20% glycerol after carefully removing the meninges and the pial vessels, embedded in gelatin or agarose (15% gelatin or 3% agarose) hardened at 4°C over night creating a 5–10mm-thick mantle surrounding the brain, further hardened in 4% formaldehyde at 4°C for up to 3 days, deeply frozen at -60°C and serially sectioned using a

cryomicrotome (Jung, Nussloch, Germany). One brain (C7) was embedded in celloidin as recently described [54] and was cut into serial 440-500µm-thick coronal sections using a sliding microtome (Polycut, Cambridge Instruments, UK). From each hemisphere, every second or third section was stained with gallocyanin (a Nissl stain) as previously described [51]. Fixation and tissue processing were performed at the Morphological Brain Research Unit, University of Wuerzburg (Germany) under identical conditions for all brains (except for brain C7 that was embedded in celloidin instead of gelatin).

Stereological analyses

Stereological analyses were performed with a stereology workstation equipped with a Zeiss Axioplan II microscope (Carl Zeiss MicroImaging, Thornwood, NY, USA), Plan-Neofluar objectives 2.5x (N.A. = 0.075) and 40x (N.A. = 1.30), Fluar objectives $10x$ (N.A. = 0.5) and 20x (N.A. = 0.75), a Microfire CCD camera (Optronics, Goleta, CA, USA), a motorized stage (Ludl Electronics, Hawthorne, NY, USA), and stereology software (StereoInvestigator, Version 10, MBF Bioscience, Williston, VT, USA).

The left hemispheres were selected and the regions of interest were delineated at low magnification (using the 2.5x objective) according to established criteria in the literature [19]. The regions of interest (ROI) were layers II-III and layers V-VI of the dACC region, corresponding to Brodmann's area 24, subregions 24a', 24b' and 24c' as described by Vogt [19]. Grossly, the most anterior section started from the crossing of the corpus callosum, and extended approximately 2 cm posteriorly to the border with area 23. Every 6th section was sampled in a consistent and unbiased manner, generating a serial section series of 5–7 sections within our ROIs. Actual section thickness after histological processing was determined as described in [55].

The volumes of the dACC region (layers II-III and layers V-VI) were analyzed using the Cavalieri principle [56]. Briefly, the area of the delineated ROIs was multiplied by the serial section interval and the actual measured section thickness. Estimation population of total neuron and total glial cell (astrocytes and oligodendrocytes) numbers in layers II-III and layers V-VI separately was evaluated using the Optical Fractionator method, see [56]. Starting with a random section number, a systematic random sampling throughout the ROI was performed. Pilot experiments were conducted to select appropriate counting frames and grid sizes, generating approximately 500 counted objects (either neurons or glia) in a total series, consistently generating coefficient of errors (CE) <0.1, see [56]. Details of the final counting procedures are summarized in Table 2. Neurons were identified by their characteristic morphology with dark and recognizable nucleoli, a large soma with a distinct cell nucleus surrounded by fainter cytoplasm, and dendrites emerging from the soma. Glial cells were identified by their smaller size, lack of processes, and dark homogenous staining. Astrocytes are nearly indistinguishable from oligodendrocytes in gallocyanin-stained materials [51], and were pooled together as glial cells. The neuron and glial cell densities were calculated as the ratio of the total cell number (either neuron or glia) and the Cavalieri volume of the ROIs. The glia/neuron index (GNI) was calculated as the ratio of the glial cell density to the neuron density.

Statistical analysis

For both patients with schizophrenia and controls, means and standard error of the mean (SEM) were calculated for age at death, age at onset of illness, post-mortem interval (PMI), and fixation time (Table 1). Means and SEM were also calculated (separately for patients with schizophrenia and controls) for the analyzed variables, dACC volume, total neurons, total glia, neuron density, glial density and GNI (separately for layers II–III and V-VI). Comparisons between patients with schizophrenia and controls were performed using

generalized linear model multivariate analysis (MANOVA), with diagnosis as fixed factor and the following variables as covariates: post-mortem interval (PMI), fixation time, age of controls or adjusted illness duration (AID) of patients with schizophrenia (AID is calculated as individual age at death minus age at onset plus the mean age at onset of all patients with schizophrenia). Note that use of the actual individual illness duration of the patients with schizophrenia instead of the adjusted ones as covariate would have caused invalid results of the MANOVA model because there was no illness duration of the controls. The mean illness duration of the patients with schizophrenia was significantly different from the mean age of the controls (two-tailed Student's *t*-test; $p < 0.001$) whereas the mean adjusted illness duration was not ($p = 0.974$). For each investigated variable, the two investigated regions of interest (layers II-III and V-VI) were tested simultaneously. In all analyses an effect was considered statistically significant if its associated p -value was <0.05. Calculations were performed using SPSS (Version 18.0.1 for Mac, PASW, Chicago, IL, USA). Graphical analysis was done with GraphPad Prism (Version 4 for Macintosh; GraphPad Software, La Jolla, CA, USA).

Photography

Photomicrographs of full brain sections were scanned at high resolution (600 dpi) using an Epson Perfection 2450 Photo scanner (Figure 2A). Photomicrographs of magnified regions were produced by digital photography using the stereology workstation with the StereoInvestigator software described above. Figures 2B and 3CD were generated using the Virtual Slide module, using either a 2.5x objective and a grid size of approximately 5x6 (Figure 2B), or a 10x objective and a grid size of 1x4 (Figure 3CD), grid size adjusted to the size of the area acquired. Figure 3AB was generated using the Deep Focus module with the 40x objective, acquiring images every 3–5µm through a section thickness of 20–30µm. All editing was performed using Adobe Photoshop CS4.

RESULTS

The region of interest

The region of the dACC used in our study encompassed the areas 24a'b'c', as delineated by Vogt and coworkers [19, 30], corresponding functionally to the cognitive subdivision of the ACC, as discussed by Bush et al. [31] (Figure 1). A representative coronal section through the dACC is shown in Figure 2. The investigated ROI extended from the rostralmost to the caudalmost levels of areas 24a', b', and c' (supplementary Figures 1–2). The ROI was also subdivided into superficial (layers II-III) and deep (layers V-VI) cortical layers because of known differences in cortical layer development and connectivity patterns [57]. The cytoarchitecture and the cellular features of the representative area 24b' is shown in Figure 3 (and supplementary Figure 3).

In human post-mortem Nissl-stained materials, oligodendrocytes may be difficult to distinguish reliably from astrocytes. However, it is known that astrocytosis (which is indicative of glial scarring and a gliotic reaction) is not present in schizophrenia [58], thus changes in glial cell numbers would suggest underlying changes in oligodendrocyte numbers. For histological analyses, no histochemical or immunohistochemical marker for specific for oligodendrocytes is sufficiently specific and reliable for quantification purposes on human postmortem tissue. As a consequence, in our Nissl-stained materials, we counted astrocytes and oligodendrocytes together as glial cells.

Observations in controls versus patients with schizophrenia

In the dorsal anterior cingulate region, we analyzed the dACC volume, the estimated population of total number of neurons, the estimated population of total number of glial

cells, the estimated neuron density, the estimated glial cell density, and the glia/neuron index (GNI), in layers II-III and layers V-VI of area 24a'b'c'. These variables were analyzed between groups (controls versus patients with schizophrenia) and against age and adjusted illness duration.

Using rigorous anatomical and stereological criteria, we found no differences between groups in either cortical volume, total neuron numbers, or total number of glial cells in the dACC (supplementary Figure 4). Similarly, we found no differences between groups in neuron density, glial density, or GNI values (Figure 4), as well as between groups when adjusting for illness duration, showing (not shown). These findings of an apparent intact dACC cytoarchitecture in schizophrenia stand in contrast to other observations of decreases as well as increases in neuronal densities in the cingulate or prefrontal cortex in patients with schizophrenia (see Discussion). Glial cell densities appeared to decrease with age and illness duration in both superficial and deep layers. However, as white matter volume generates an inverse U-shaped developmental pattern in healthy patients [59], regression analysis was not used to assess this apparent decrease (see Discussion).

Observations in superficial versus deeper cortical layers

In contrast to the lack of differences between groups, we found anticipated differences between cortical layers in both patients with schizophrenia and in controls (Figure 4). Overall, the neuron densities were higher in superficial layers, whereas the glial densities were higher in the deep layers. This was also reflected in the calculation of the glia/neuron ratios, the GNI in layers II-III was approximately 1.5 in both groups, and the GNI in layers V-VI was approximately 2.3 in both groups. We found differences in the superficial layers versus the deep layers in glial densities in the dACC, in both controls and patients with schizophrenia, reflecting that there were more glial cells in the deep layers.

DISCUSSION

Using rigorous anatomical and stereological criteria, we observed no changes in neuronal or glial cell densities in the dACC in patients with schizophrenia compared to controls from a cohort of age- and gender-matched subjects, spanning from 20 to70 years of age. Neither did we detect any changes in the volume, total neuron numbers, or total glial cell numbers in the dACC between patients with schizophrenia and controls. These findings do not support previous observations of apparent changes in neuron densities in prefrontal and cingulate regions in patients with schizophrenia. We did however find expected differences between superficial and deep layers in both neuron and glial cell densities, as well as the GNI, in the dACC in both controls and patients with schizophrenia. These observations can be attributed the importance of subregion- and layer- specificity when addressing neuropathological changes in schizophrenia. Differences in histopathological changes in post-mortem human tissue may reflect several methodological and technical differences (such as sample size, age-matched cases, left and right hemispheres, stereological versus two-dimensional analyses, section thickness, staining procedures, regional delineation, etc.). The present results emphasize the necessity to provide meticulously detailed methodological and analytical information in quantitative neuropathological studies.

Previous morphological data on cortical thickness and imaging data on regional volumes have suggested that different areas in schizophrenia may have a reduced volume [41, 44, 45, 60–63]. Despite the abundance of imaging data suggesting reduced volume of various cortical regions, these findings have been difficult to correlate with post-mortem histopathological data. Both subregion delineation as well as analytical differences between imaging and histopathological data may lie behind these difficulties. When investigating particular subregions of the anterior cingulate and prefrontal cortex, for example, Koo and

colleagues found volume differences in the subgenual and rostral ACC, but no volume differences in the dorsal ACC or in the posterior cingulate cortex [45]. A study by Narr and colleagues [43] found decreases in ACC thickness in women, whereas in men only slight decreases in dACC thickness in the right hemisphere were observed, with an overall increase in prefrontal cortical thickness. In histological materials, the reduction in brain volume in select areas has primarily been attributed to a reduction in neuronal size and in reduced neuropil, rather than a loss of neurons [64, 65]. Others have found indications of a decreased cortical thickness and smaller pyramidal soma size in area 24abc as well as in area 25 in schizophrenia, but with no changes in neuronal densities [66].

Studies on neuronal densities in the prefrontal cortex and the anterior cingulate cortex have led to inconclusive outcomes because rigorous stereological approaches have not been employed, different Brodmann areas have been pooled, or generalized theories have been presented from the results of only a few cortical regions. For these reasons, some authors find no changes in neuronal densities in schizophrenia, some find decreases in neuronal density, while others found increases in neuronal density. For methodological purposes in quantitative neuroanatomy, we discuss some examples of these inconclusive findings below.

Pakkenberg and coworkers employed stereology but found no changes in neuronal densities, yet these studies pooled several Brodmann areas from the prefrontal and cingulate cortex [67, 68]. In contrast, Stark and colleagues found no differences in neuronal densities in area 24 or 32 [36], and Bouras and coworkers found no differences in neuronal densities in area 24abc [66]. Benes and coworkers employed a 2D counting approach and found lower densities of neurons in the cingulate cortex while subregions were not specified [69, 70]. These reductions in neuron densities in the cingulate have not been replicated by others using stereological methods. Selemon and coworkers employed stereology and found higher neuronal densities in the prefrontal cortex, yet mainly focused on area 9 and 46 [39, 40]. They also found smaller soma sizes [71] and smaller frontal grey matter volume [60] and extrapolated these results to a generalized view of changes in the prefrontal cortex in schizophrenia [64]. In the ACC, Chana and colleagues also found increased neuronal densities and smaller soma sizes using stereologicalal methods [72]. Despite these discrepancies, these findings have supported the altered circuitry hypothesis of schizophrenia [64, 73]. However, the difficulty in comparing and evaluating these reports remains, as they contain methodological limitations that may have had influence the outcome.

Not as many studies have investigated glial density changes in schizophrenia. This may be due to the fact that there were apparently no astrogliotic changes occurring in schizophrenia [58], and the influence of potential oligodendrocyte changes was perhaps overlooked. In the prefrontal and cingulate regions, Hof and colleagues found decreased oligodendrocyte densities in area 9 [35], and Stark and coworkers found decreased oligodendocyte densities in area 24 but not in area 32 in schizophrenia [36]. In comparison, Segal and coworkers found no differences in oligodendrocytes densities in the cingulum bundle [37]. Reductions in glial densities have also been found in the subgenual anterior cingulate cortex in depression [74, 75]. These observations have previously suggested that changes in oligodendrocyte numbers may be indicative of aberrant wiring and conduction abnormalities in schizophrenia.

In assessments of neuropathological changes in cortical and subcortical architecture and histology, crucial components are to specify the region and subregions studied, since subtle changes may be present within one region [19,30, 31, 76]. For example, using the same cases as in this study, Kreczmanski et al. [46] found reduced volume and total neuron numbers, but not densities, in the putamen and lateral nucleus of the amygdala, but not in

other striatal or amygdala subregions. Within the ACC, the pregenual and subgenual regions have been shown to be relevant to emotional functions [77] and are strongly connected with the amygdala [78], and have primarily been implicated in depression [34, 79]. However the rACC has mainly been studied with imaging, as reliable morphometric or stereological investigations of the pregenual cortex are notoriously difficult in view of its curved morphology. Lastly, layer specificity in schizophrenia studies is particularly important considering that schizophrenia is also considered to be a neurodevelopmental disorder [86, 87, 88]. In this context, it is noteworthy that deep layers myelinate before superficial layers during development [80]. This may be the reason for higher numbers of oligodendrocytes in the deep layers, as found in this study. Any aberrations in cortical layering may be of substantial importance for the development of schizophrenia.

The theories on the aetiology and psychopathology of schizophrenia are multifaceted, from neurochemical to wiring and development. The fact that a myelination deficit potentially represents a substrate for the clinical symptoms seen in patients with schizophrenia merits to be reviewed in the context of developmental aberrations [81, 82] and altered cortical circuitry [65, 75, 83–86]. The myelin theory on schizophrenia originated from the finding of myelin pathology and downregulation of the expression of myelin-related genes in schizophrenia [1]. Myelin and oligodendrocyte integrity are required for adequate functioning of neurons and hence of circuitry. It has been shown that both white matter volume and grey matter volume changes during development and aging [59]. Sowell and coworkers found that the white matter development followed and inverted U-pattern, with white matter volume increasing until midlife and then declining, whereas the grey matter volume progressively decreases with age [59]. Therefore, a myelination deficit in schizophrenia, which has gained much support in recent years from white matter architecture and tractography imaging studies [85, 87–90], may be directly linked to both the theories of development and circuitry.

Our results, however, suggest that any neuropathological and neurohistological changes are highly region- and layer-specific. Myelin changes previously observed may perhaps be specifically localized to other areas of the prefrontal cortex, such as area 9 [35]. This is an intriguing outcome that points to fact that the dACC may not undergo as severe changes in its cellular makeup in schizophrenia as has been generally believed, and stresses the need for further detailed postmortem investigations, at the level of identified and specific domains of grey and white matter.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

Malin Höistad and Patrick Hof designed the experiments and wrote the paper. Malin Höistad conducted all of the stereological analyses, acquired the images, and performed all of the data analyses. Helmut Heinsen acquired, sectioned and stained all brains and coordinated all communication with clinical and pathological departments. Christoph Schmitz verified all statistics, stereological design, and helped with the study design. Bridget Wicinski helped with the stereology and acquisition of images, and maintained the collection of histological materials. This study was supported by NIH grant MH066392 (PRH), grants from Stanley Medical Research Institute (#02R-258 and #04R-674 to HH, CS, and PRH), and by a postdoctoral fellowship from the Swedish Brain Foundation to Malin Höistad. The authors are deeply indebted to the patients and their families who have made this study possible.

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Figure 1.

(A) The anterior cingulate region (ACC) as delineated by Vogt, encompassing the rostral areas 24a-c and the dorsal areas 24a'-c', bordering the paracingulate area 32 and 32' and the subgenual area 25. Modified from [19]. (B) The rostral (blue/affective) and dorsal (red/ cognitive) anterior cingulate regions depicted on a reconstructed MRI, of the medial surface of the right hemisphere of a human brain. Modified from [31].

Figure 2.

Inset: A mid-sagittal photograph of the left hemisphere of a human brain showing the cingulate gyrus. The vertical line corresponds to a representative coronal section that contains the region of interest. Scale $bar = 1$ cm. (A) Gallocyanine-stained coronal section showing the localization of the investigated region, the dorsal anterior cingulate (dACC, within dashed box) in a control subject. Scale bars $= 1$ cm. (B) Photomontage of the dACC corresponding to areas 24a'-c', illustrating the cytoarchitectonic features at each level from a control subject. Arrowheads indicate borders among areas 33, 24a', 24b', 24c', and 32', respectively, moving dorsally from the corpus callosum. Scale bar = 1 mm.

Figure 3.

Photomicrographs depicting typical cellular features in layer V of area 24b' in a control subject (A) and in a patient with schizophrenia (B). Note the many satellite oligodendrocytes surrounding pyramidal cell perikarya. Scale bar = 25 µm. Photomicrographs depicting the six cortical layers in area 24b' in a control (C) and in a patient with schizophrenia (D). Scale $bar = 100 \mu m$.

Figure 4.

Figure 4. (A-D) Variations in neurons in controls (open circles) and in patients with schizophrenia (filled circles), in superficial (II-III) and deep (V-VI) layers of the dACC volume (corresponding to area 24a'-c'). Estimated neuron densities between groups (A, C), and across age (B, D).

(E-H) Variations in glial cells (oligodendrocytes and astrocytes) in controls (open circles) and in patients with schizophrenia (filled circles), in superficial (II-III) and deep (V-VI) layers of the dACC volume (corresponding to area 24a'-c'). Estimated glial densities between groups (E, G), and across age (F, H).

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(I-L) Variations in GNI ratios (glial density/neuron density) in controls (open circles) and in patients with schizophrenia (filled circles), in superficial (II-III) and deep (V-VI) layers of the dACC volume (corresponding to area 24a'-c'). GNI ratios between groups (I, K), and across age (J, L).

Table 1

Clinical characteristics of cases included in the present study. Clinical characteristics of cases included in the present study.

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C - control; S - schizophrenia; C - control; S - schizophrenia; $\stackrel{*}{\wedge}$ djusted illness duration, see Materials and Methods. Adjusted illness duration, see Materials and Methods.

 $t_{\text{Case S1 and S2 had long post-mortem intervals, however they were both found within 1h of death and were kept at 4°C until autopsisy.}$ $t_{\text{Case S1 and S2 had long post-mortern intervals, however they were both found within 1h of death and were kept at 4°C until autopsisy.}$

DSM-IV: Diagnostic and Statistical Manual of Mental Disorders, 4th Edition [49]. DSM-IV: Diagnostic and Statistical Manual of Mental Disorders, 4th Edition [49]. ICD-10: International Statistical Classification of Diseases and Related Health Problems [50]. ICD-10: International Statistical Classification of Diseases and Related Health Problems [50].

 V ICD-10

Table 2

Details of the stereological parameters.

