Dopamine D2 Receptor Levels in Striatum, Thalamus, Substantia Nigra, Limbic Regions, and Cortex in Schizophrenic Subjects

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Abstract

Background—Studies in schizophrenics have reported dopaminergic abnormalities in striatum, substantia nigra, thalamus, anterior cingulate, hippocampus, and cortex which have been related to positive symptoms and cognitive impairments.

Methods—[18F]fallypride PET studies were performed in off medication or never medicated schizophrenic subjects [N = 11, 6 M, 5 F; mean age of 30.5 ± 8.0 (S.D.); 4 drug naive] and age matched healthy subjects [N = 11, 5M, 6F, mean age of 31.6 ± 9.2 (S.D.)] to examine dopamine D2 receptor (DA D2r) levels in the caudate, putamen, ventral striatum, medial thalamus, posterior thalamus, substantia nigra, amygdala, temporal cortex, anterior cingulate, and hippocampus.

Results—In schizophrenic subjects increased DA D2r levels were seen in the substantia nigra bilaterally; decreased levels were seen in the left medial thalamus. Correlations of symptoms with region of interest data demonstrated a significant correlation of disorganized thinking/nonparanoid delusions with the right temporal cortex region of interest (r = 0.94, P = 0.0001) which remained significant after correction for multiple comparisons (P<0.03). Correlations of symptoms with parametric images of DA D2r levels revealed no significant clusters of correlations with negative symptoms, but significant clusters of positive correlations of total positive symptoms, delusions and bizarre behavior with the lateral and anterior temporal cortex, and hallucinations with the left ventral striatum.
Conclusions—The results of this study demonstrate abnormal DA D_2r mediated neurotransmission in the substantia nigra consistent with nigral dysfunction in schizophrenia and suggest that both temporal cortical and ventral striatal DA D_2r mediate positive symptoms.

Keywords
Dopamine D_2 receptors; Schizophrenia; fallypride; substantia nigra; thalamus; delusions; hallucinations

Abnormal dopaminergic neurotransmission has been implicated in the positive symptoms and cognitive deficits seen in schizophrenia (1-5). Recent studies suggest abnormal function of GABAergic/glutamatergic cortical microcircuits in schizophrenia resulting in dysfunction of cortical pyramidal glutamatergic neurons (6) which provide a major excitatory afferent projection to the substantia nigra (7). Dysfunction of this projection results in nigral dysfunction and increased striatal DA release (8,9, and 10) which has been positively correlated with positive symptoms (11). Prefrontal cortical glutamatergic afferents to the ventral tegmental area (VTA) synapse directly on mesocortical DA neurons; it has been hypothesized that dysfunction of this projection leads to decreased cortical DA release(12) which is believed to be a factor in the cognitive impairments seen in schizophrenia(4). As dopamine D_2 receptors (DA D_2r) directly modulate cortical GABAergic interneurons which modulate cortical pyramidal neurons (13,14), ventral midbrain DA neurons (15,16), and DA release in striatal and extrastriatal regions (17), DA D_2r are of considerable interest in schizophrenia.

Consistent with the hypothesis of decreased cortical DA release in schizophrenia, post mortem studies have reported decreased dopaminergic innervation in medial temporal cortex, dorsolateral prefrontal cortex and hippocampus (18-20), and DOPAC levels in the anterior cingulate (21). Some imaging studies of extrastriatal DA D_2r in schizophrenic subjects have reported decreased DA D_2r levels in the anterior cingulate and temporal cortex, but most have not (22-27); the most frequent finding is increased medial thalamic DA D_2r levels (24,25, 26). While there have been variable results, a recent study of DA D_1r in schizophrenic subjects reported increased frontal cortical levels which were negatively correlated with performance on a working memory task (28-30), The increased DA D_1r levels were interpreted as being consistent with decreased frontal cortical DA levels. Post mortem studies of dopaminergic function in the the substantia nigra in schizophrenic subjects have reported increased levels of tyrosine hydroxylase (31), tyrosine hydroxylase mRNA (32), homovanillic acid (31), and DA D_2r (33) consistent with nigral dysfunction. Imaging studies have largely failed to examine substantia nigra DA D_2r. Both post mortem and imaging studies have reported increased striatal DA synthesis, DA levels, and DA release which has been correlated with positive symptoms (34-41). In contrast to post mortem studies which have reported increased striatal DA D_2r levels (42,43), most imaging but not all studies of striatal DA D_2r have reported unaltered levels in schizophrenia (44,45,46). However, one imaging study of striatal DA D_2r performed before and after DA depletion with alphamethylparatyrosine demonstrated normal levels prior to DA depletion but increased DA D_2r levels after depletion consistent with both increased striatal DA release and increased total DA D_2r levels(36). The discrepancy between post mortem studies and imaging studies with benzamide radioligands may be due to the occupancy of striatal DA D_2r by increased levels of extracellular DA. Overall, the available post mortem and imaging data are consistent with the hypothesis of decreased cortical DA release, nigral DA neuronal dysfunction and increased striatal DA release in schizophrenia.

Previous imaging studies of DA D_2r in medication free schizophrenic subjects have evaluated either striatal or extrastriatal DA D_2r levels (22-27,44,45). In the current study, positron emission tomography (PET) with [18F]fallypride was utilized. [18F]Fallypride is a very high affinity, specific benzamide PET radioligand for the DA D2 receptor (K_D = 0.03nM) and is
the only currently available radioligand which allows estimation of both striatal and extrastriatal DA D2r levels (22,24,27,44-48). Given the hypothesis of nigral dysfunction in schizophrenia(1,8-10,12), the lack of previous imaging studies of the substantia nigra DA D2r, post mortem findings consistent with nigral dysfunction(31-33), and the ability of PET [18F]fallypride studies to estimate nigral DA D2r levels(48,49), we specifically examined this region. Other regions previously reported to have altered DA D2r levels, the anterior cingulate, temporal cortex, and medial thalamus(22-26), were examined. As significant correlations of symptoms with regional DA D2r levels have been reported (22-27), correlations of positive and negative symptoms with regional DA D2r levels were assessed.

Methods

Subjects

This study was conducted under protocols approved by the Vanderbilt University and Centerstone Mental Health Center Institutional Review Boards. All subjects were judged capable of giving informed consent by a senior research psychiatrist, and provided informed consent for this study. Subjects meeting the DSM IV criteria (American Psychiatric Association, 1994), and Research Diagnostic Criteria (50) for the diagnosis of schizophrenia between the ages of 18 and 45 were recruited. The diagnosis of schizophrenia was established by the Structured Clinical Interview for DSM IV Axis I disorders (51) and checklist. Schizophrenic subjects [N = 11, 6 M, 5 F; mean age of 30.5 ± 8.0 (S.D.) years and age range of 20–45 years] were either never treated (N=4) or were off medication for at least three weeks (Table 1). The BPRS (6 item scales), SAPS and SANS were administered to each subject; mean total BPRS, SAPS and SANS scores were 28.8 ± 7.0 (S.D.), 9.8 ± 3.1 (S.D.), and 9.4 ± 4.0 (S.D.) respectively. Age matched healthy subjects [N = 11, 5M, 6F, mean age of 31.6 ± 9.2 (S.D.) years and age range of 21–45 years] were recruited as well. Significant medical conditions, and previous or current substance abuse were exclusion criteria for all subjects.

Data Acquisition and Analysis

MRI scans of the brain were performed using a G.E. 1.5T Signa LXi MRI scanner. High resolution T1-weighted gradient echo acquisitions in the sagittal plane (1.2 -1.3 mm thick slices) and coronal planes (1.4–1.5 mm thick slices), axial spin density weighted and T2-weighted (3 mm thick slices) acquisitions were obtained. PET scans were performed using a GE Advance PET scanner in the 3-D acquisition mode. [18F]Fallypride (4–5 mCi, specific activity >2,000 Ci/mmol, maximum mass dose of less than 2.5 nanomoles) was injected intravenously over a 20 second period; serial scans of increasing duration were obtained for 210 minutes, allowing stable estimates of binding potentials in all regions (47-49). A measured attenuation correction was utilized.

Serial PET scans were coregistered to each other and to thin section T1 weighted MRI images using a rigid body, mutual information algorithm (52,53), and reoriented to the ACPC line. Regions of interest were identified on thin section T1 weighted MRI images, and transferred to coregistered PET studies. The putamen, and caudate were manually drawn by a neuroradiologist (RMK) on axial slices from 2 to 12 mm above the ACPC line. The ventral striatum was defined using the criteria of Mawlawi (54). Sobel filtering was performed on high resolution gradient echo MRI images of the brain (55), but did not provide reliable boundaries for delineation of the dorsomedial thalamus and pulvinar. We used anatomic landmarks to delineate the medial thalamus and posterior thalamus, which approximated the boundaries of the dorsomedial thalamus and pulvinar (56). The medial thalamus was delineated on slices from 2 to 12 above the ACPC line; the posterior border was the coronal plane of the posterior commissure, the medial border the midline, the anterior boundary the foramen of Monro and the lateral border extended up to 1 cm from the midline. The anterior border of the posterior...
thalamus was the coronal plane of the posterior commissure, the medial and posterior borders
the edge of the thalamus as it projects into the quadrigeminal plate cistern, and the lateral border
the posterior limb of the internal capsule. The substantia nigra/VTA is located in the ventral
midbrain 9-14 mm below the ACPC line (56) and can be readily visualized in the midbrain on
PET [18F]fallypride scans (57). Substantia nigra regions of interest were manually drawn to
adjust for inter-individual variability by a neuroradiologist (RMK); the intersubject coefficient
of variation for the substantia nigra region was 8.7 % (57). The amygdala can be visualized in
the on MRI scans just anterior to the tip of the temporal horn of the lateral ventricle and deep
to the uncus (57); the amygdala is located 6 to 20 mm below the ACPC line, 12 to 28 mm
lateral to the midline, and from 2 to 12 mm behind the plane of the anterior commissure (56).
To decrease partial voluming from the striatum, amygdala regions of interest were drawn on
MRI images from 10 to 16 mm below the plane of the ACPC. Temporal cortical regions of
interest were manually drawn on axial MRI images from 35 to 25 mm below the ACPC. Our
previous studies have shown excellent inter-subject reliability for these regions of interest, i.e.
inter-subject coefficients of variation of 6.8 to 15.9 percent (57). The anterior cingulate was
delineated as extending from superior to the axial plane through the ACPC in the pregenual
region superiorly and posteriorly to the coronal plane through the anterior commissure. The
hippocampus was manually delineated on the coronal MRI images from the tip of the temporal
horn anteriorly to the last coronal slice in which it would be reliably identified posteriorly.
Regional DA D2 receptors were estimated using the reference region method with a cerebellar
reference region (58). The cerebellum is an appropriate reference region as less than 3 percent
of cerebellar uptake is specific binding to DA D2 receptors and reference region method
estimates of binding potentials are highly correlated (r>0.99) with modeled estimates using a
metabolite corrected plasma input function (47,59-61). Parametric images of DA D2 were
coregistered across subjects using an elastic deformation algorithm (62).

Statistical Analysis

Region of interest data were analyzed using a repeated measures MANOVA with region and
hemisphere as within subject factors, and group (schizophrenic, healthy control) as a between
subject factor with age as a covariate. Definition of the hippocampal region of interest was
problematic in one subject, and this subject was left out of the analysis. When the MANOVA
was performed with this subject but without the hippocampus as a region, no conclusion was
changed. Because age has a significant effect on DA D2 receptors, independent group two-tailed
t-tests covaried for age were used to test for group differences in regional binding potentials.
To evaluate positive symptoms, the total SAPS scores, and global SAPS scores for
hallucinations, delusions and bizarre behavior as well the BPRS positive symptom score, and
BPRS scores for suspiciousness (Item 11), hallucinations (Item 12), and disorganized thinking/
nonparanoid delusions (Item 15) were measured. Negative symptoms were examined using the
SANS. Correlations of symptom scores with regions of interest were performed using a
Pearson product moment correlation and significance evaluated using a two-tailed t-test.
Bonferroni correction was used to correct for multiple comparisons. Correlations of symptom
scores with parametric DA D2 images were calculated on a voxel basis using a Pearson product
moment correlation and significance evaluated using a two-tailed t-test. To assess the
significance of clusters of significant correlations, corrections for multiple comparisons were
made using the method of Forman as implemented in the AFNI program (63). The critical
threshold for the voxelwise analysis was P<0.01 and a minimum cluster size of 24 voxels.
These clusters were significant at P<0.001 corrected for multiple comparisons.

Results

A repeated measures MANOVA was performed with region and hemisphere as within subject
factors, group (schizophrenic, healthy control) as a between subject factors, and with age as a
covariate. No significant main effect of hemisphere, no group × hemisphere interaction, or group × hemisphere × region interaction was seen. There was a significant effect of region, \( F(7,13) = 36.78, p<0.00001 \), reflecting the large differences in regional binding potentials (Table 2). There was no main effect of group. However, there was a significant region × group interaction, \( F(7,13) = 6.00, p<0.005 \). There was also a region × age interaction, \( F(7,13) = 4.60, p<0.010 \) reflecting decreases in regional binding potentials with age which were greater for cortical than subcortical regions. To explore the regions responsible for the significant region × group interaction, independent group two tailed t-tests covaried for age were performed to examine which region(s) might account for this interaction. These tests demonstrated significantly increased DA D_2_r levels in the substantia nigra/VTA bilaterally and decreased levels in the left medial thalamus (Table 2). No other region of interest demonstrated a significant difference in DA D_2_r levels between schizophrenic and healthy subjects.

Correlation of region of interest data with SAPS scores, SANS scores, BPRS positive symptom score, and BPRS scores for suspiciousness (Item 11), hallucinations (Item 12), and disorganized thinking/nonparanoid delusions (Item 15) demonstrated one correlation which survived Bonferroni correction and a second trend level correlation. DA D_2_r levels in the right temporal cortex region of interest were positively correlated with the BPRS score for disorganized thinking/nonparanoid delusions, \( r = 0.94, P = 0.0001 \) uncorrected for multiple comparisons, \( P<0.03 \) after correction for multiple comparisons. The left temporal cortical region of interest demonstrated a trend level correlation with the BPRS score for disorganized thinking/nonparanoid delusions after corrections for multiple comparisons, \( r = 0.92, P = 0.0003, \) uncorrected for multiple comparisons, \( P<0.08, \) corrected for multiple comparisons.

Correlations of symptoms with regional DA D_2_r levels performed using voxelwise analysis revealed no significant clusters of correlations of regional DA D_2_r levels with either the total SANS score or individual SANS scores. Significant clusters of highly positive correlations of regional DA D_2_r levels were seen with the total SAPS score, and global SANS scores for delusions, hallucinations and bizarre behavior. Two clusters of highly positive correlations (146 voxels on the right, mean \( r = 0.85 \); 131 voxels on the left, mean \( r = 0.86 \)) were seen with total SANS scores; these clusters involved the posterior portions of the inferior, middle and superior temporal gyri and extended superiorly into the supramarginal gyrus of the parietal lobe in both cerebral hemispheres (Figure 1). Significant clusters of correlations of SAPS global delusions scores with DA D_2_r levels was seen in the lateral aspects of the right and left anterior temporal cortex extending into the temporal tips laterally (50 voxels on the right, mean \( r = 0.84 \); 80 voxels on the left, mean \( r = 0.86 \)) (Figure 2). Similarly, correlations of the BPRS score for disorganized thinking/nonparanoid delusions (Item 15) with DA D_2 receptor levels demonstrated a similar cluster of positive correlations in the left anterior temporal cortex (80 voxels, mean \( r = 0.85 \)). The SANS global bizarre behavior scores demonstrated bilateral clusters of positive correlations (184 voxels on the right, mean \( r = 0.85 \); 179 voxels on the left, mean \( r = 0.84 \)). The cluster on the left involved the posterior portions of the inferior and superior temporal gyri as well the mid to posterior portions of the middle temporal gyrus with extension into the inferior parietal lobule. The cluster on the right also involved the inferior, middle, and superior temporal gyri and inferior parietal lobule; it extended further anteriorly in the sulcus between the superior and middle temporal gyrus, but had less extension into the posterior superior temporal gyrus and inferior parietal lobule than the cluster on the left (Figure 3). In contrast, the SAPS global scores for hallucinations demonstrated positive correlations with the left ventral striatum (31 voxels, mean \( r = 0.84 \)) but not with cortical regions (Figure 4). BPRS scores for hallucinations demonstrated a similar left ventral striatal cluster of positive correlations (41 voxels, mean \( r = 0.87 \)).
Discussion

The results of this study indicate that there are increased DA D_2r levels in the substantia nigra/VTA and decreased DA D_2r levels in the left medial thalamus. The increased levels of nigral/VTA DA D_2r seen in the current study are consistent with the one post mortem study of nigral DA D_2r in schizophrenics which also reported increased levels (33). DA D_2r in the substantia nigra are largely inhibitory autoreceptors on nigral DA neurons (15,16). As discussed above, post mortem studies have also reported increased nigral levels of tyrosine hydroxylase, tyrosine hydroxylase mRNA, and homovanillic acid (31,32) in the substantia nigra of schizophrenic subjects. The findings in both the current study and previous post mortem studies demonstrate both increased inhibitory nigral DA D_2 autoreceptor levels and increased DA synthesis and release suggesting dysregulation of midbrain dopaminergic neurons in schizophrenic subjects. Similar findings, i.e. increased total DA D_2r levels and increased DA synthesis and release (34-43), have been reported in the striatum of schizophrenic subjects and suggest that similar dysregulation of dopaminergic neurotransmission occurs in both nigra and striatum. The factor(s) responsible for increased nigral and striatal DA D_2r levels when increased extracellular DA levels are preset are unclear.

The VTA, dorsal tier of the zona compacta of the substantia nigra and retrorubal fields provide dopaminergic innervation to limbic and cortical regions and so are of considerable interest in schizophrenia (64,65). The resolution of the PET scanner used in this study is insufficient to distinguish changes in these areas from the ventral tier of the zona compacta which provides dopaminergic innervation to the striatum. While the PET scanner used in this study does not have sufficient resolution to provide complete quantitative recovery of DA D_2r levels in the substantia nigra, published calculated estimates of quantitative recovery for the substantia nigra indicate that the the 5-6 mm resolution of the scanner does allow substantial recovery of quantitation (66). Consistent with these calculations are studies which indicate the ability of the scanner used to estimate SN DA D_2r occupancies by a number of antipsychotic drugs as well as the changes in apparent SN DA D_2r levels following DA release and DA depletion (49,57,59,67). There has been one recent [123I]epidepride SPECT study which has reported decreased levels of midbrain uptake in schizophrenic subjects (68). The low resolution of SPECT relative to the size of the substantia nigra does not allow separation of nigral DA D_2r from those in other structures. In addition, the lack of a scatter correction, the use of a ratio method using cerebellum as a reference region prior to the attainment of a transient equilibrium, and the variability in this ratio due to lipophilic metabolites of [123I]epidepride in the cerebellum makes interpretation of these results difficult (69-71).

The results of this study confirm the previously reported finding of decreased left medial thalamic DA D_2 rceptor levels in schizophrenic subjects (24-26). An autoradiographic study of human thalamic DA D_2r has reported a heterogenous and nuclear specific distribution of DA D_2r with highest levels in the midline and intralaminar nuclei of the thalamus; levels in the dorsomedial nucleus were at least two fold lower than in the midline and intralaminar nuclei (72). While the dorsomedial nucleus accounts for most of the medial thalamic region of interest, the midline and intralaminar nuclei are included in this region of interest. As a number of cognitive functions and behaviors which are impaired in schizophrenia are mediated by prefrontal cortical/basal ganglia/medial thalamic circuits (73), a loss of DA D_2r in the dorsomedial nucleus of the thalamus may contribute to these impairments. The thalamic intralaminar nuclei project to frontal cortex, striatum and limbic regions providing feedback from the thalamus to these regions (74,75); this feedback is affected by DA D_2r in these nuclei providing an additional site for modulation of prefrontal cortical/basal ganglia/medial thalamic circuit function. The apparent reduction in medial thalamic DA D_2r levels may reflect loss of medial thalamic neurons expressing DA D_2r consistent with imaging and post mortem studies reporting decreased medial thalamic volume and neuronal numbers (55,76-79), a loss of
autoreceptors on medial thalamic dopaminergic projections, or an increase in thalamic DA release (80). However, increases in thalamic DA release are unlikely to cause the decrease in apparent left medial thalamic DA D2 receptor levels as d-amphetamine administration produces only a 3% decline in medial thalamic [18F]fallypride binding potentials in humans (57).

The current results suggest that different positive symptoms are mediated by DA D2 in different regions. Scores for delusions and bizarre behavior are positively correlated with anterior temporal/temporal tip, and lateral temporal/inferior parietal cortical DA D2, respectively, while hallucinations are positively correlated with left ventral striatal but not cortical DA D2. Consistent with these correlations are cerebral blood flow studies in schizophrenic subjects which found positive correlations of left ventral striatal and left temporal tip blood flow in schizophrenic subjects with a reality distortion factor principally related to hallucinations and delusions (81). A comprehensive review of neuropathological lesions producing schizophrenic symptoms reported an association of striatal lesions with auditory hallucinations, whereas left temporal lobe lesions were associated with delusions (82). The differences in regional correlations for hallucinations and delusions raise the possibility that hallucinations and delusions may be differentially affected by antipsychotic drugs which produce preferential occupancy of temporal cortical versus striatal DA D2 (60, 67, 83, 84). The lack of significant clusters of correlations with negative symptoms suggests that these symptoms may not be mediated by DA D2 neurotransmission.

The positive correlations of positive symptoms with cortical DA D2 levels are similar to a recent [123I]epidepride SPECT study which reported a positive correlation of positive symptoms with frontal cortical DA D2 levels in males but not females (27). In subjects with bipolar disorder psychosis has been correlated with increased striatal DA D2 (85) consistent with the results of the current study. Although previous studies, (23-26) have reported negative correlations of medial, lateral and/or total thalamic DA D2 receptor levels with positive symptoms as measured by the BPRS or with the PANSS general psychopathological scores, no significant correlations of symptom scores with medial thalamic regions of interest were seen.

There a number of potential limitations in this study. These include the small number of subjects studied and the fact that seven of the eleven subjects studied had received previous neuroleptic treatment. While the number of schizophrenic subjects examined in the current study is similar to other PET studies of extrastriatal DA D2 which have studied 7 to 15 subjects, a larger cohort may provide more reliable estimates of DA D2 levels in schizophrenia (22-26). The largest study of extrastriatal DA D2 in unmedicated schizophrenics, a SPECT study of 25 subjects, did not evaluate the regions found to be abnormal in the current study (27). As increased, decreased, and unchanged levels of DA D2 are seen in the current study, it is unlikely that the increased levels seen reflect receptor upregulation due to previous therapy or the decreased levels residual antipsychotic drug effects. Although subjects were not carefully matched for smoking status, it is unlikely that the current results were affected by smoking status as extrastriatal DA D2 levels are not affected by smoking status (86). While females were not carefully matched for menstrual status, one study of the effect of menstrual status on DA D2 levels in humans (86) found no statistically significant effect while a second, older study (87) reported a small effect but no statistical significance was reported; the effect, if any, is small. Finally, extracellular DA levels may be altered in schizophrenia and affect the apparent levels of DA D2 as [18F]fallypride has been shown to be sensitive to extracellular DA levels (49, 57).

In conclusion, the results of this study demonstrate increased substantia nigra and decreased left medial thalamic DA D2 levels in off medication schizophrenic subjects. Positive
correlations of positive symptoms with temporal cortical and left ventral striatal striatal DA D2r levels were found. The increased substantia nigra DA D2r levels are consistent with the hypothesized nigral dysfunction in schizophrenia. The positive correlations of hallucinations with ventral striatal DA D2r levels, and delusions and bizarre behavior with temporal cortical receptor levels provides additional evidence for the role of DA D2r mediated neurotransmission in these key psychotic symptoms, and suggests that these symptoms may be mediated by DA D2r in different brain regions.

Acknowledgments

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Figure 1.
Sagittal (A,B), axial (C), and coronal (D) images through significant clusters of correlations of total SAPS scores with regional DA D_{2r} levels. Two significant clusters are seen involving the posterior portions of the inferior, middle and superior temporal gyri with extension superiorly into the supramarginal gyrus of the parietal lobe in both cerebral hemispheres. The cluster on the right (146 voxels, mean r =0.85) was similar in size to the cluster on the left (131 voxels, mean r=0.86)
Figure 2.
Two significant clusters of correlations of the SAPS global score for delusions with DA D_{2r} levels are seen in the right and left anterolateral temporal cortex extending into the temporal tips. The cluster on the left (80 voxels on the left, mean r=0.86) is larger than the cluster on the right (50 voxels on the right, mean r=0.84).
Figure 3.
Sagittal left (A) and right (B), axial (C), and coronal (D) images through significant clusters of correlations of SAPS global scores for bizarre behavior with regional DA D\textsubscript{2r} levels. Two significant clusters of highly positive correlations (184 voxels on the right, mean $r=0.85$; 179 voxels on the left, mean $r=0.84$) involve the mid to posterior lateral aspects of the temporal lobes with extension into the inferior parietal lobule.

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Figure 4.
A significant cluster of correlations (31 voxels, mean $r = 0.84$) of the SAPS global score for hallucinations with DA $D_2R$ levels is seen in the left ventral striatum. No other significant clusters of correlations were seen.
Table 1

Demographic data for OFF Medication Schizophrenic Subjects

<table>
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<tr>
<th>Subject Number</th>
<th>Sex</th>
<th>Age</th>
<th>Medication Free period</th>
<th>Previous medications</th>
<th>BPRS Score (6 item scales)</th>
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<tr>
<td>1</td>
<td>F</td>
<td>36</td>
<td>3 weeks</td>
<td>quetiapine</td>
<td>30</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>32</td>
<td>9 weeks</td>
<td>quetiapine</td>
<td>32</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>22</td>
<td>5 weeks</td>
<td>olanzapine</td>
<td>38</td>
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<tr>
<td>4</td>
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<tr>
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<td>39</td>
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<td>19</td>
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<td>11</td>
<td>M</td>
<td>26</td>
<td>3 weeks</td>
<td>olanzapine</td>
<td>22</td>
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Table 2

Binding potentials for regions of interest sampled in 11 unmedicated schizophrenic subjects and 11 age matched healthy subjects. Significance level was estimated using an independent group, two tailed t-test covered for age.

<table>
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<th>Region</th>
<th>Schizophrenic</th>
<th>Normal</th>
<th>Significance level</th>
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<td></td>
<td>R</td>
<td>L</td>
<td>R</td>
</tr>
<tr>
<td>Medial Thalamus</td>
<td>4.21±0.58</td>
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<td>4.30±0.62</td>
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<td>Substantia Nigra</td>
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<td>2.44±0.22</td>
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<td>Hippocampus</td>
<td>1.57±0.41</td>
<td>1.68±0.38</td>
<td>1.48±0.26</td>
</tr>
<tr>
<td>Temporal Cortex</td>
<td>1.52±0.33</td>
<td>1.63±0.33</td>
<td>1.59±0.18</td>
</tr>
<tr>
<td>Caudate</td>
<td>30.70±3.30</td>
<td>31.81±4.12</td>
<td>32.25±2.08</td>
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<tr>
<td>Putamen</td>
<td>36.52±4.36</td>
<td>35.00±4.46</td>
<td>37.02±2.56</td>
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<tr>
<td>Ventral Striatum</td>
<td>18.80±3.66</td>
<td>19.49±3.49</td>
<td>18.10±2.81</td>
</tr>
<tr>
<td>Amygdala</td>
<td>3.28±0.43</td>
<td>3.20±0.61</td>
<td>3.23±0.27</td>
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</tbody>
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