Tract-based Analysis of Magnetization Transfer Ratio and Diffusion Tensor Imaging of the Frontal and Frontotemporal Connections in Schizophrenia

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Background: In the pathophysiology of schizophrenia, aberrant connectivity between brain regions may be a central feature. Diffusion tensor imaging (DTI) studies have shown altered fractional anisotropy (FA) in white brain matter in schizophrenia. Focal reductions in myelin have been suggested in patients using magnetization transfer ratio (MTR) imaging but to what extent schizophrenia may be related to changes in MTR measured along entire fiber bundles is still unknown. Methods: DTI and MTR images were acquired with a 1.5-T scanner in 40 schizophrenia patients and compared with those of 40 healthy participants. The mean FA and mean MTR were measured along the genu of the corpus callosum and the left and right uncinate fasciculus. Results: A higher mean MTR of 1% was found in the right uncinate fasciculus in patients compared with healthy participants. A significant negative correlation between age and mean FA in the left uncinate fasciculus was found in schizophrenia patients but not in healthy participants. Conclusions: Decreased FA in the left uncinate fasciculus may be more prominent in patients with longer illness duration. The increased mean MTR in the right uncinate fasciculus could reflect a compensatory role for myelin in these fibers or possibly represent aberrant frontotemporal connectivity.

Key words: schizophrenia/diffusion tensor imaging/magnetization transfer ratio/tract-based analysis/uncinate fasciculus/genu of the corpus callosum

Introduction

Using standard anatomical magnetic resonance imaging (MRI) methods, numerous studies have revealed that gray matter volume decreases in the brains of patients with schizophrenia, in particular in the frontotemporal regions. These regions do not operate in isolation but form large-scale neural networks for which normal interregional communication is a prerequisite for proper functioning. Indeed, from about the time schizophrenia was defined, it was suggested that aberrant connectivity between these brain regions is a central feature of the disease. Recently, evidence is accumulating that the integrity of white matter fibers connecting the spatially distinct gray matter regions is compromised in schizophrenia and that impaired functioning of white matter is part of its pathophysiology.

Diffusion tensor imaging (DTI) and magnetization transfer ratio (MTR) imaging are MRI methods that allow us to study different aspects of the connecting white matter fiber bundles in vivo that cannot be measured using standard T1- or T2-weighted MRI. Using DTI, decreases in fractional anisotropy (FA)—which reflects microstructural directionality and to a certain extent fiber integrity—have been found in several studies in schizophrenia (for reviews, see Kanaan et al, Kubicki et al, and Konrad et al). Most studies applied voxel-based analysis of DTI images. More recently, fiber-tracking techniques are being used to infer fiber integrity along complete tracts. Using such tract-based analysis methods, characteristic values such as FA can be measured along fiber tracts and subsequently their average values between groups can be compared. As such, tract-based analysis methods are optimal to detect subtle group-related changes that may occur along entire fiber tracts. This is in contrast to voxel-based analyses, which are optimized to detect focal differences in white matter.

In the few studies that applied a tract-based analysis to measure FA in schizophrenia, no differences in mean FA values between groups were found. However, a significant correlation between higher age and lower mean FA was shown in schizophrenia patients and not in healthy participants. This group-by-age interaction suggests that the deficits found in schizophrenia may be more prominent in patients with longer illness duration. Indeed, recently, reductions in FA were found in
chronically ill and not in first-episode schizophrenia patients compared with healthy participants.17

MTR imaging measures the presence of macromolecules in tissue, including myelin, and thus may provide additional information to DTI on white matter integrity.18–21 A few MTR imaging studies of white matter have been completed in schizophrenia. In the first study, using an region of interest (ROI) approach, reduced mean MTR values were found in the left and right temporal regions but not in the frontal, parietal, and occipital regions.22 Similarly, reductions in mean MTR values were found in the splenium but not in the genu of the corpus callosum.23 Bilaterally reduced MTR was found using a voxel-based analysis in the uncinate fasciculus (with left reductions greater than right reductions) in first-episode schizophrenia patients,24 although another voxel-based analysis study did not find significant reductions in MTR.25 In the one combined DTI and MTR voxel-based analysis study in schizophrenia, decreased MTR was found in the posterior cingulum bundle, corpus callosum, fornix, right internal capsule, and superior occipitofrontal fasciculus bilaterally in schizophrenia patients compared with healthy participants.26 Tract-based analysis of FA in combination with MTR may provide more information about to what extent possible group differences in FA can be attributed to differences in myelination. To our knowledge, no tract-based analysis has been done for MTR in schizophrenia and its associations with age.

In this study, we measured fiber integrity and myelin concentration over entire fiber tracts in schizophrenia. We compared both FA and MTR along averaged white matter tracts27 computed for the uncinate fasciculi, which connect the temporal and frontal lobes, and the genu of the corpus callosum (genu), which connects the frontal lobes of both hemispheres. We selected fibers connecting the frontal lobes of the left and right hemispheres and fibers connecting the frontal and temporal lobes because these brain areas are known to be involved in the pathophysiology of schizophrenia. Indeed, numerous studies have reported decreases in frontal and temporal gray matter volumes and densities.28–30 Moreover, earlier we found decreases in white matter density in the anterior corpus callosum using a voxel-based morphometry approach. Therefore, we hypothesized that fiber integrity as measured using DTI and MTR along the uncinate fasciculi and anterior corpus callosum would be compromised in patients with schizophrenia as compared with healthy participants. In addition, we investigated associations of FA and MTR with age, severity of positive and negative symptoms, outcome, age at onset, duration of illness, and antipsychotic medication intake.

Methods

Subjects

Forty patients with schizophrenia and 40 healthy participants matched for age, gender, handedness, and parental education participated in this study. The healthy participants were recruited by means of local newspaper advertisements. All subjects participated after written informed consent was obtained. The study was approved by the medical ethics committee for research in humans (medisch ethische toetsingscommissie) of the University Medical Center Utrecht, The Netherlands.

All participants underwent extensive psychiatric assessment procedures using the Comprehensive Assessment of Symptoms and History (CASH). Patients met Diagnostic and Statistical Manual of Mental Disorders (Fourth Edition) criteria for schizophrenia. “Age of onset of illness” was defined as the age at which the patients experienced psychotic symptoms for the first time, as obtained from the CASH interview, and Schedule for Affective Disorder and Schizophrenia Lifetime version assessed by 2 independent raters. Diagnostic consensus was achieved in the presence of a psychiatrist. “Duration of illness” was defined as the time between the age of onset of illness and the age at the time of the MRI scan. All healthy participants met research diagnostic criteria for “never mentally ill” and had no first-degree family member with a mental illness or second-degree relatives with a psychotic disorder.

All patients were receiving typical, or atypical, antipsychotic medication at the time of the scan. A table from the Dutch National Health Service (Commissie Farmaceutische Hulp van het College voor Zorgverzekeringen, 2002) was used to calculate the cumulative dosage of typical antipsychotics during the scan interval and to derive the haloperidol equivalents. For atypical antipsychotics, the respective pharmaceutical companies suggested how to convert dosage into haloperidol equivalents. For atypical antipsychotics, the respective pharmaceutical companies suggested how to convert dosage into haloperidol equivalents (clozapine, 40:1; olanzapine, 2.5:1; risperidone, 1:1, sulpiride, 170:1;quetiapine, 50:1; and sertindole, 2:1). Drug use was assessed with the Composite International Diagnostic Interview. Four patients and 1 healthy participant met the criteria for drug abuse, and 1 patient met the criteria for drug dependency. Drugs used included cannabis (in all 6 subjects) and others (3). For demographics, see table 1.

Image Acquisition

MRI scans of the whole brain were made on a 1.5-T Intera Achieva Philips System at the University Medical Center Utrecht using a 6-element SENSE receiver head coil. For each subject DTI scans, an MTR scan and a high-resolution T1-weighted scan (used for anatomical reference) were collected. First, a 3-dimensional T1-weighted coronal (spooled gradient) echo scan of the whole head was acquired (256 × 256 matrix; echo time (TE) = 4.6 ms; repetition time (TR) = 30 ms; flip angle = 30 degrees; 160–180 contiguous slices; total scan duration = 405–456 s; 1 × 1 × 1.2 mm³ voxels; field of view (FOV) = 256 mm/70%; and parallel imaging applied in both phase-encoding directions with SENSE factor = 1.5). For white matter fiber tract reconstruction and computation of the FA
Table 1. Demographic Data

<table>
<thead>
<tr>
<th></th>
<th>Patients With Schizophrenia (n = 40)</th>
<th>Healthy Participants (n = 40)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male/female, no.</td>
<td>30/10</td>
<td>29/11</td>
</tr>
<tr>
<td>Age, y°</td>
<td>26.8 (5.8)</td>
<td>28.0 (7.7)</td>
</tr>
<tr>
<td>Age, y range</td>
<td>20–41</td>
<td>18–45</td>
</tr>
<tr>
<td>Height, cm°</td>
<td>179.3 (9.2)</td>
<td>182.5 (9.0)</td>
</tr>
<tr>
<td>Height, cm range</td>
<td>163–198</td>
<td>167–204</td>
</tr>
<tr>
<td>Weight, kg°</td>
<td>76.3 (13.3)</td>
<td>74.3 (9.1)</td>
</tr>
<tr>
<td>Weight, kg range</td>
<td>55–110</td>
<td>57–92</td>
</tr>
<tr>
<td>Handedness—right/left/ambidexter, no.</td>
<td>37/3/0</td>
<td>35/5/0</td>
</tr>
<tr>
<td>Duration of illness</td>
<td>25.1 (17.4)</td>
<td></td>
</tr>
<tr>
<td>Medication at the time of the scan&lt;bc</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Typical antipsychotics, no.</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Median (range)</td>
<td>3.5 (4.7)</td>
<td></td>
</tr>
<tr>
<td>Haldol equation</td>
<td>Atypical antipsychotics, no.</td>
<td></td>
</tr>
<tr>
<td>Median (range)</td>
<td>7.1 (12)</td>
<td></td>
</tr>
<tr>
<td>Haldol equation</td>
<td>No medication at the time of the scan, no.</td>
<td>0</td>
</tr>
<tr>
<td>Cumulative medication&lt;ad</td>
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<td></td>
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<tr>
<td>Typical antipsychotics, no.</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>106.5</td>
<td></td>
</tr>
<tr>
<td>Atypical antipsychotics, no.</td>
<td>22</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>6208.4 (6830.9)</td>
<td></td>
</tr>
<tr>
<td>Typical + atypical antipsychotics, no.</td>
<td>13</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>10956.2 (9640.3)</td>
<td></td>
</tr>
<tr>
<td>PANSS positive symptoms</td>
<td>15.7 (5.6)</td>
<td></td>
</tr>
<tr>
<td>PANSS negative symptoms</td>
<td>15.6 (5.7)</td>
<td></td>
</tr>
<tr>
<td>PANSS general symptoms</td>
<td>31.0 (7.2)</td>
<td></td>
</tr>
<tr>
<td>PANSS total score</td>
<td>62.3 (15.8)</td>
<td></td>
</tr>
<tr>
<td>CAN total score</td>
<td>14.4 (7.9)</td>
<td></td>
</tr>
</tbody>
</table>

a P < .01.
bValues are mean ± SD.
cIn 7 patients medication information was unavailable.
dHaloperidol, broomperidol, pipamperon, thioridazine, flufenazine, perphenazine, zuclopenthixol, penfluridol, pimozide, fluspirilene, largactil, and flufenazine were considered typical antipsychotic drugs; clozapine, risperidone, olanzapine, and sertindole were considered atypical antipsychotic drugs.

dIn 4 patients cumulative medication use was unavailable.

The 2 DTI scans were simultaneously realigned and corrected for possible gradient-induced distortions. A robust estimation of the diffusion tensors was obtained using M-estimators to limit the influence of possible outliers. From the diffusion tensors the FA was computed. The FA represents the elongation of the diffusion profile and assumes values between 0 (isotropic diffusion) and 1 (pure 1-dimensional diffusion) and is defined by

\[
FA = \sqrt{\frac{3}{2} \left( \frac{(\lambda_1 - \lambda)^2 + (\lambda_2 - \lambda)^2 + (\lambda_3 - \lambda)^2}{\lambda_1 + \lambda_2 + \lambda_3} \right)},
\]

where \(\lambda_1\) represents the eigenvalue of the diffusion tensor’s major eigenvector pointing in the direction parallel to the fiber bundle and \(\lambda_2\) and \(\lambda_3\) represent the eigenvalues of both minor eigenvectors pointing in the radial directions.

To compute the MTR, the second volume of the magnetization transfer scan was rigidly aligned with the first volume using the ANIMAL software package. Mutual information was used as a similarity metric. The MTR was computed on the image \(I_m\) with the magnetization pre-pulse and the image \(I_0\) without the magnetization pre-pulse using the equation \(\text{MTR}=\frac{(I_0 - I_m)}{I_0} \times 100\). The MTR is expressed as a percentage where 0% represents no signal reduction and 100% represents total signal reduction due to magnetization transfer.

For both the MTR (here the scan without the magnetization pre-pulse was used) and the T1-weighted scan, the rigid transformations were determined that spatially aligned them with the diffusion-unweighted \((b = 0 \text{ s/mm}^2)\) volume of the DTI scan using mutual information as similarity metric. For each subject, a nonlinear transformation was computed using the ANIMAL software package that spatially aligns the subject’s T1-weighted scan with a T1-weighted model brain. This nonlinear transformation was used at a later stage to warp the reconstructed tracts into the model space.
Fiber Tracking and Fiber Bundle Selection

A multiple ROI fiber bundle selection approach was used to select the fiber tracts of interest where the reconstruction of the tracts was performed in native space and the selection and analysis of the tracts representing the genu and the left and right uncinate fasciculi were performed in model space.

In the first step, all possible tracts in the brain were reconstructed individually in native space using the diffusion tensor images with an in-house implementation of the fiber assignment by continuous tensor fusion images with an in-house implementation reconstructed individually in native space using the diffusion tensor images with an in-house implementation.

In the first step, all possible tracts in the brain were reconstructed individually in native space using the diffusion tensor images with an in-house implementation of the fiber assignment by continuous tensor fusion images with an in-house implementation reconstructed individually in native space using the diffusion tensor images with an in-house implementation.

In the second step, the ROIs that were needed to select the tracts representing the genu and the left and right uncinate fasciculi were manually delineated on the model brain using an average map with color-coded directional information of the main direction of the tensors (figure 1a). For the left and right uncinate fasciculi, the ROIs were placed in the temporal and prefrontal regions (figure 1b). For the genu, 2 ROIs were placed, 1 in the left and 1 in the right prefrontal region (figure 1c).

In addition to remove spurious tracts, a third ROI was placed at the midline, anterior to the ventricles (the blue colored ROI in figure 1c).

Instead of transforming the ROIs defined in model space to each individual subject (native space) in order to select the genu and uncinate fasciuli tracts from the reconstructed tracts, we transformed all reconstructed tracts from each subject into model space. The advantage is that tracts are geometric representations (poly-lines) that can be transformed without the need for interpolation. This is not the case when the ROIs are transformed from the model to native space because they are defined by voxels.

Computation of Average Fibers

We computed an average fiber (in model space) of the genu and left and right uncinate fasciculi for each individual subject. Figure 1 shows an example of the computation of the average fiber for the right uncinate fasciculus.

In short, first a spline representation of the shape of the original reconstructed fiber tracts (figure 1d) was created. This spline was then divided in 2-mm regular intervals starting from the center of the spline. This center was defined by the geometric center of the midpoint coordinates of all tracts of the fiber bundle perpendicularly projected onto the spline (the center is denoted by the red plane in figure 1e). For each of these points, we defined planes that are perpendicular to the spline. These planes (the position and direction of these planes are denoted by the green disks in figure 1e) were used to resample the original reconstructed fiber tracts (figure 1d). Each plane was cross-sectioned with the original tracts of the fiber bundle. The cross-sectional coordinates were averaged to form the coordinate of the average fiber in this plane. For each separate plane, the MTR and FA values of the cross-sectional coordinates of the original tracts were averaged to produce the average FA and MTR values for that plane. The average cross-sectional coordinates of all the planes together with the corresponding average FA and MTR values then formed the average fiber. For each subject, the final mean FA and mean MTR values were computed by averaging the FA and MTR values, respectively, of all points of an average fiber. These final mean FA and MTR values were computed for each subject.

### Table 2. Fractional Anisotropy (FA) and Magnetization Transfer Ratio (MTR) Values in Individual Tracts of Patients With Schizophrenia and Healthy Participants

<table>
<thead>
<tr>
<th>Tracts</th>
<th>Mean (SD) FA Values</th>
<th>Mean (SD) MTR Values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Patients</td>
<td>Healthy Participants</td>
</tr>
<tr>
<td>Genu of corpus callosum</td>
<td>0.58 (0.04)</td>
<td>0.57 (0.04)</td>
</tr>
<tr>
<td>Left uncinate fasciculus</td>
<td>0.46 (0.03)</td>
<td>0.47 (0.03)</td>
</tr>
<tr>
<td>Right uncinate fasciculus</td>
<td>0.47 (0.03)</td>
<td>0.46 (0.03)</td>
</tr>
</tbody>
</table>

*P < .0001.

*aTract-based comparisons between groups are corrected for age and gender. FA and MTR measurements are based on 39 patients and 40 control subjects for the right uncinate fasciculus because in 1 patient the right uncinate fasciculus could not be tracked.

*bAfter removal of 1 outlier from healthy participants (MTR values including outlier were 61.28 (0.90) for healthy participants and F(1,76) was 0.51).
and for each average fiber representation of the genu and the left and right uncinate fasciculus.

In addition, we computed group average fibers over all individual average fibers for visualization purposes (figure 2). These group average fibers allowed us to resample each individual average fiber in order to visualize the locality of possible group effects.

Statistical Analysis

Mean MTR and mean FA measurements were analyzed using SPSS 15.0. Data were examined for outliers, extreme values, and normality of distribution. No normality transformations were needed. Using general linear modeling, multiple univariate analyses of covariance were done with the mean MTR and mean FA of each separate average fiber as dependent variables, group (schizophrenia, healthy participants) entering as fixed factor, and age and gender as covariates. For dependent variables that differed significantly between groups, Pearson product-moment correlations were computed for each group separately (or in patients only) between mean MTR and mean FA, between mean MTR/FA and level of education, and between mean MTR/FA and clinical variables. For the genu and the left and right uncinate fasciculus, the correlations between age and mean FA and age and mean MTR were computed separately for schizophrenia patients and healthy participants. A z statistic was computed (after Fisher’s Z transform) to test if these correlations differ between groups.

Results

FA and MTR

A significant increase in mean MTR was found in the right uncinate fasciculus in patients with schizophrenia compared to healthy participants ($F(1,76) = 15.56, P = .0001$), which remained significant after Bonferroni correction for multiple comparisons (table 2). No changes in mean FA were found in the right uncinate fasciculus in patients with schizophrenia compared with healthy counterparts.
participants ($F(1,76) = 0.76, P = .31$). No significant changes in mean FA or mean MTR in the left uncinate fasciculus or genu were found between groups. Excluding subjects with drug abuse or drug dependency from the sample did not alter the findings. To obtain additional information on whether the measured MTR group differences could be related to myelin concentrations, a post hoc analysis was performed on the so-called transverse diffusivity. The transverse diffusivity is defined as the average of the 2 minor eigenvalues, $(\lambda_2 + \lambda_3)/2$, and shows a higher correlation with myelin concentrations than FA.\(^{38}\) No significant group differences were found in transverse diffusivity for any of the fiber bundles.

A significant difference between correlations of mean FA with age ($z = -2.28, P = .02$) was found between schizophrenia patients and healthy participants in the left uncinate fasciculus (figure 3). To determine whether possible medication effects could explain this finding, an additional post hoc analysis was performed where the cumulative medication was regressed out from the measured FA values in patients before computing the difference in correlations between age and FA between patients with schizophrenia and healthy comparison subjects. This resulted in a difference in correlations between age and FA in patients with schizophrenia compared with healthy participants for the left uncinate fasciculus that was significant at the trend

Fig. 2. Fractional Anisotropy (FA) and Magnetization Transfer Ratio (MTR) Measurements along the Uncinate Fasciculi. (a) FA and MTR values in patients (red rectangle up) and healthy participants (blue rectangle down) with a 2-mm interval along the left uncinate fasciculus (left) and right uncinate fasciculus (right). Note that the bars represent +1 and −1 standard error of the mean. For both uncinate fasciculi, no differences in FA were found between patients and healthy participants. The MTR in the right uncinate fasciculus measured along the complete fiber was significantly higher in patients than in healthy participants. In contrast, comparison at a local level (ie, per 2-mm interval) did not reveal significant differences in MTR or FA (after Bonferroni correction for multiple comparisons with the number of comparisons set to 43). (b) Local differences in MTR between patients and healthy participants along the average group fiber of the right uncinate fasciculus (more reddish color represents higher local MTR in patients).
level ($z = -1.91, P = .056$). For the genu, the difference ($z = -1.43$) found between schizophrenia patients and healthy participants did not reach significance.

Correlation analysis between mean MTR and mean FA in the genu and the left and right uncinate fasciculus in patients or in healthy participants did not reveal any significant correlations.

**Associations of the Uncinate Fasciculus with Clinical Symptoms, Education, and Antipsychotic Medication**

No significant correlations of mean FA and mean MTR in the right uncinate fasciculus were found with age at first symptoms, duration of illness, positive and negative and general symptoms of the positive and negative syndrome scale, total Camberwell Assessment of Need score, and antipsychotic medication intake at the time of the scan. There were no significant correlations (corrected for age, gender, and duration of illness) between cumulative antipsychotic medication use and mean MTR or between cumulative antipsychotic medication use and mean FA, in the genu and the left or right uncinate fasciculus.

A significant correlation was found between mean MTR in the left uncinate fasciculus with the extent of negative symptoms ($r = -0.55, P < .0001$), which remained significant after Bonferroni correction for multiple comparisons. There were no significant correlations of mean MTR and mean FA in the left uncinate fasciculus with any of the other clinical variables. No significant correlations were found between mean MTR in the genu and mean MTR in the right uncinate fasciculus with any of the clinical variables.

**Discussion**

In this study, FA and the magnetization transfer ratio (MTR) were measured using DTI and MTR imaging along the uncinate fasciculi and the genu of the corpus callosum in the brains of 40 patients with schizophrenia and compared with those of 40 healthy participants. Our main finding is a significant higher mean MTR of 1%, with no differences in mean FA, in the right uncinate fasciculus in the patients with schizophrenia compared with healthy participants. The MTR increases were found along large parts of the fiber (figure 2) and were not limited to local changes of the tract which serves as an indication that these increases reflect schizophrenia-related changes that affect entire fiber tracts. In addition, a significant negative correlation between age and mean FA in the left uncinate fasciculus was found in schizophrenia patients but not in healthy participants.

The significant difference in correlations between age and FA in patients compared with healthy participants (figure 3) suggests that in patients FA reductions become more apparent with age (mean age in patients was 26.8 years, range between 20 and 41 years). However, medication effects could be an important factor that should be taken into account when interpreting the results. The results of the additional post hoc analysis where cumulative medication was taken into account indicate that, although medication effects may play a role, they cannot completely explain the measured differences in correlations between age and FA values. Our results support recent findings of a tract-based analysis that reported significant differences in correlations between age and FA in patients with schizophrenia compared with healthy participants for the left and right uncinate fasciculus combined. The fact that we found this correlation in the left and not in the right uncinate fasciculus is in keeping with the previous findings showing decreases in FA in the left but not in the right uncinate fasciculus of schizophrenia patients who were on average 43 years.

If the difference in correlation between age and FA predominantly relates to changes in myelination, then one may expect to find similar differences in correlation between age and MTR because MTR appears to show a higher correlation with myelin levels than FA. We did not find a difference in correlations between age and MTR in patients compared with healthy participants. Therefore, our findings at least suggest that the

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Fig. 3. Correlation Between Age and Fractional Anisotropy for Patients ($r = -0.36, P = .02$) (Red Rectangle Up) and Healthy Participants ($r = -0.15, P = .36$) (Blue Rectangle Down) in the Left Uncinate Fasciculus (Left) and Right Uncinate Fasciculus (Right).
Our finding of increased MTR in the right uncinate fasciculus suggests altered connectivity between the right frontal and temporal cortices in schizophrenia and may imply increased myelin concentrations in the largest fiber bundle connecting the medial frontal and temporal cortices. Because increased myelin concentrations along the fiber have been associated with more efficient axonal relaxation, an increase in myelin in the uncinate fasciculus suggests an increased level of signal transfer between frontal and temporal cortices that could reflect a compensation mechanism for decreased interhemispheric connectivity in schizophrenia. The notion that increased MTR reflects increased myelination in the right hemisphere is in keeping with the results of functional studies showing increased levels of activation in the right frontal cortex in schizophrenia. In this context, it should be noted that increased communication speed does not necessarily lead to increased communication efficiency. The increased MTR could also result from compensation of aberrant brain functioning elsewhere in the brain, for instance in the left hemisphere. If, however, increases in myelin are responsible for the measured increase in MTR, then one might also expect (to some extent) a reduction in transverse diffusivity, which is defined by the average of the minor eigenvalues of the diffusion tensor. However, the post hoc analysis of the transverse diffusivity in the right uncinate fasciculus did not reveal any group-related differences, suggesting that our MTR findings may not (only) reflect increased myelination. An alternative explanation for the increased mean MTR in the right uncinate fasciculus in patients should therefore be considered, namely that it might reflect compromised connectivity not directly related to myelin.

MTR does not only depend on the macromolecule concentration but also (among others) depend on T1 relaxation times of free water molecules. Prolonged T1 relaxation times in white matter do not reflect a difference in myelination, but a change in the free/bound water fraction. Interestingly, prolonged T1 relaxation times in the white matter of the right hemisphere have been reported for schizophrenia. Such an increase in T1 could effectively lead to an increase in MTR. An alternative explanation for the increased mean MTR in the right uncinate fasciculus in patients might involve changes in metabolites in white matter. Although we can only speculate, the increased mean MTR may be (indirectly) related to increased glutamate and glutamine levels. In that case, the increased mean MTR could be due to a prolonged T1 relaxation time reflecting an altered free/bound water fraction possibly related to changes in glial glutamate uptake. MTR levels have been negatively correlated with glutamate and glutamine levels in white matter as measured with MR spectroscopy.

The average FA in the genu of the corpus callosum appeared lower in older patients as compared with older control participants, but this finding did not reach significance. In the genu of the corpus callosum, a decrease in FA in patients compared with healthy participants was reported earlier, although others did not find group differences in mean FA.

The absence of such a difference in FA in this study may be due to the relative low mean age (22.6 years) combined with an age span of 18–45 years of our population because such FA reductions appear to become more pronounced with age. Another possible explanation may be that the reported differences in FA are of more focal nature while tract-based analysis is most sensitive to group differences that are found along large parts of the tracts. Both explanations may hold for the fact that in a previous study from our group structural decreases in density were found using voxel-based morphometry in the genu for patients with schizophrenia (age span 16.3–67.9 years, mean age 36.6 years).

Although the measured difference in MTR in the right uncinate fasciculus was only in the order of 1%, we note that it is a robust and statistically highly significant finding because increased MTR values were consistently found in schizophrenia along large parts of the tract. Although small, these differences could, for instance, have large consequences for the synchronization of the signal because small alterations of signal transport along large parts of the fiber may cumulate into large alterations of various aspects of the signal transport (eg, signaling speed and signal response function).

A limitation of this study is that the applied fiber-tracking algorithm requires sufficient directional information to successfully reconstruct the fibers. If, at a certain point, this information is not available (for instance, due to crossings with other fibers), then the algorithm cannot reconstruct the complete fiber tract. Another limitation is that like DTI, MTR is an indirect measure of white matter microstructure. Therefore one should interpret group differences between FA or MTR values with caution as both FA and MTR are indirect measures and other factors then fiber orientation or macromolecule content may alter FA or MTR values as well.

In conclusion, a subtle but significant increase in mean MTR was found along large parts of the right uncinate fasciculus of schizophrenia patients. This increase appears to be specific for the right uncinate fasciculus and points toward increased connectivity between the medial frontal cortex and temporal pole provided that increases in myelin fully account for the increases found in mean MTR. Possibly, the right uncinate fasciculus may be part of a compensation mechanism for aberrant functioning of the left uncinate fasciculus, as suggested by the decreased FA in the older patients with schizophrenia. However, other mechanisms such as prolonged T1 relaxation times may also be implicated.
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