White matter integrity in schizophrenia and bipolar disorder: Tract- and voxel-based analyses of diffusion data from the Connectom scanner

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ARTICLE INFO

Keywords:
Schizophrenia
Bipolar Disorder
DTI
TRACULA
TBSS

ABSTRACT

Background: Diffusion imaging abnormalities have been associated with schizophrenia (SZ) and bipolar disorder (BD), indicating impaired structural connectivity. Newer methods permit the automated reconstruction of major white matter tracts from diffusion-weighted MR images in each individual’s native space. Using high-definition diffusion data from SZ and BP subjects, we investigated brain white matter integrity using both an automated tract-based and voxel-based methods.

Methods: Using a protocol matched to the NIH (Young-Adult) Human Connectome Project (and collected on the same customized ‘Connectom’ scanner), diffusion scans were acquired from 87 total participants (aged 18–30), grouped as SZ (n = 24), BD (n = 33) and healthy controls (n = 30). Fractional anisotropy (FA) of eighteen white matter tracts were analyzed using the TRACULA software. Voxel-wise statistical analyses of diffusion data was carried out using the tract-based spatial statistics (TBSS) software. TRACULA group effects and clinical correlations were investigated using analyses of variance and multiple regression.

Results: TRACULA analysis identified a trend towards lower tract FA in SZ patients, most significantly in the left anterior thalamic radiation (ATR; p = .04). TBSS results showed significantly lower FA voxels bilaterally within the cerebellum and unilaterally within the left ATR, posterior thalamic radiation, corticospinal tract, and superior longitudinal fasciculus in SZ patients compared to controls (FDR corrected p < .05). FA in BD patients did not significantly differ from controls using either TRACULA or TBSS. Multiple regression showed FA of the ATR as predicting chronic mania (p = .0005) and the cingulum-angular bundle as predicting recent mania (p = .02) in patients. TBSS showed chronic mania correlating with FA voxels within the left ATR and corpus callosum.

Conclusions: White matter abnormality in SZ varies in severity across different white matter tract regions. Our results indicate that voxel-based analysis of diffusion data is more sensitive than tract-based analysis in identifying such abnormalities. Absence of white matter abnormality in BD may be related to medication effects and age.

1. Introduction

Schizophrenia (SZ) and bipolar disorder (BD) are relatively prevalent psychiatric disorders, that cause significant disability (Eaton et al., 2008). The hallmark symptoms of schizophrenia involve psychosis (i.e. delusions or hallucinations), disorganization and negative symptoms; while bipolar disorder classically consists of varying mood episodes, usually involving mania/hypomania and depression. BD shares symptomatic overlap with SZ (Coryell et al., 2001), is prevalent at increased rates in families with SZ (Kendler et al., 1993), and shares chromosomal linkage with SZ (Goes et al., 2007; Park et al., 2004; Potash et al., 2003). Additionally, at least 50% of those with BP have experienced psychosis in their lifetime (Coryell et al., 2001). While the etiology of both SZ and BP remains elusive, there is accumulating evidence that abnormalities in brain connectivity have a major role in the neurobiology of these disorders.

Diffusion imaging is a powerful non-invasive tool for examining structural brain connectivity based on patterns of water diffusion in neural tissue. With this imaging modality, the diffusion tensor model is most commonly employed, and yields the frequently used fractional anisotropy (FA) measure, which is a measure of the asymmetry of water diffusion, and indirectly indexes “neuronal integrity”, putatively reflecting both myelination and organization of the white matter tracts. Diffusion imaging studies in SZ have shown decreased FA in long-range
association tracts, including the superior longitudinal fasciculus, cingulum bundle, uncinate fasciculus, inferior longitudinal fasciculus and arcuate fasciculus; projection tracts, such as the anterior thalamic radiation and the corpus callosum (Arnedo et al., 2015; Karlsgodt, 2016; Peters et al., 2010; Wheeler and Voineskos, 2014). Diagnostic methods using tensor datasets (Mukherjee et al., 2008) and have been shown to be very sensitive to differences in processing parameters (Jones et al., 2005). Recently, TRACULA (TRActs Constrained by Underlying Anatomy), an automated probabilistic tractography toolbox within FreeSurfer (Yendiki et al., 2011), was used to segment 18 major white matter tracts (Ji et al., 2017; Sprooten et al., 2016). To our knowledge, TRACULA has not been previously used to investigate white matter tract abnormality in SZ. Studying SZ and BD subjects together using identical imaging protocols, will facilitate the understanding of differences of brain connectivity of these disorders. A potential disadvantage of a tract-based approach, such as TRACULA, is that it could dilute the identification of FA abnormalities if these are not present universally across given tracts. Thus, we hypothesize that investigations of white matter integrity would be most informative if these are investigated on the advantages of each method.

Table 1
Baseline demographic and clinical characteristics across participant groups.

| Characteristic            | CN (n = 30) | SZ (n = 24) | BP (n = 33) | F/χ² | p  |
|---------------------------|-------------|-------------|-------------|------|----|
| Age                       | 24.5 (3.0)  | 24.6 (3.6)  | 26.5 (3.0)  | 3.9  | 0.02 |
| Gender (%)                |             |             |             |      |     |
| Male                      | 15 (50.0)   | 5 (20.8)    | 18 (45.5)   | 7.2  | 0.03 |
| Female                    | 15 (50.0)   | 19 (79.2)   | 15 (54.6)   |      |     |
| Ethnicity (%)             |             |             |             | 10.0 | 0.02 |
| Asian                     | 4 (13.3)    | 1 (4.2)     | 3 (9.1)     |      |     |
| Black                     | 7 (23.3)    | 16 (66.7)   | 3 (9.1)     |      |     |
| White                     | 15 (50.0)   | 5 (16.7)    | 22 (66.7)   |      |     |
| Other                     | 1 (3.3)     | 1 (4.2)     | 1 (3.0)     |      |     |
| Handedness (%)            |             |             |             | 8.1  | 0.23 |
| Left                      | 3 (10.0)    | 2 (9.5)     | 3 (9.1)     |      |     |
| Right                     | 27 (90.0)   | 19 (90.5)   | 30 (90.9)   |      |     |
| Psychotropic medication Hx (%) |         |             |             |      |     |
| Typical neuroleptic       | 0           | 8 (33.3)    | 2 (6.1)     |      |     |
| Atypical neuroleptic      | 0           | 14 (58.3)   | 12 (36.4)   |      |     |
| Lithium                   | 0           | 6 (18.2)    | 3 (9.1)     |      |     |
| Other Mood Stabilizerb    | 0           | 4 (16.7)    | 12 (36.4)   |      |     |
| SSRI/SNRI                 | 1 (3.3)     | 6 (25.0)    | 9 (27.3)    |      |     |
| Other Antidepressantc     | 0           | 2 (8.3)     | 5 (15.2)    |      |     |
| Stimulant                 | 1 (3.3)     | 0           | 0           |      |     |
| Benzodiazepines           | 0           | 3 (12.5)    | 6 (18.2)    |      |     |
| Anticholinergic           | 0           | 4 (16.7)    | 0           |      |     |
| None                      | 29 (96.7)   | 4 (16.7)    | 3 (9.1)     |      |     |
| Duration of Illness (months) | N/A       | 72.5 (42.4) | 119.2 (69.9) |      |     |
| SAPSd                      |             |             |             |      |     |
| Positive symptoms         | 0.07 (0.4)  | 3.42 (2.8)  | 1.03 (1.8)  | 23.4 | < 0.0001 |
| Disorganization symptoms  | 0.07 (0.4)  | 1.04 (1.5)  | 1.03 (1.7)  | 5.2  | 0.007  |
| SANSd                      |             |             |             |      |     |
| Negative symptoms         | 0.7 (1.4)   | 6.25 (3.0)  | 2.88 (3.3)  | 28.4 | < 0.0001|
| WERCAPe                    |             |             |             |      |     |
| Mania                     | 4.60 (5.7)  | 15.63 (9.3) | 26.21 (5.9) | 76.1 | < 0.0001|
| Psychosis                 | 0.77 (3.2)  | 30.25 (11.2)| 10.42 (9.3) | 83.9 | < 0.0001|
| YMRS                      | 0.5 (1.1)   | 1.9 (2.7)   | 3.5 (4.3)   | 7.2  | 0.001  |
| Stress                    | 19.6 (33.5) | 32.6 (22.7)| 40.5 (33.4) | 3.6  | 0.03   |

Values are given as means (SD) or number per group (%). Results derived from results of two-way ANOVA or Chi-Square analyses.

a Refers to antidepressants other than selective serotonin reuptake inhibitors (SSRI).
b Other mood stabilizers included Tegretol/Carbamazepine, Depakote/Divalproex, Trileptal/Oxcarbazepine, Topomax/Topiramate, and Lamictal/Lamotrigine.
c Maximum possible score on the Structured Assessment of Positive Symptoms (SAPS) is 16.
d Maximum possible score on the Structured Assessment of Negative Symptoms (SANS) is 20.

* WERCAP = Washington Early Recognition Center Affectivity and Psychosis Screen. Maximum possible score on the Mania section is 49. Maximum possible score on the Psychosis section is 64.
scanner was highly modified to improve diffusion imaging, by including a Siemens SC72 gradient coil and stronger gradient power supply with maximum gradient amplitude of 100 mT/m (Sotiropoulos et al., 2013). High gradient amplitudes are beneficial for diffusion MRI (dMRI) and increase the signal-to-noise ratio (SNR) over conventional MRI systems (Sotiropoulos et al., 2013). We hypothesize that our optimized dMRI protocol will be sensitive to small group differences in white matter integrity. Both groups are hypothesized to show a pattern of decreased FA in multiple tracts, with greater impairment in the schizophrenia group. TBSS is expected to identify the regions with the greatest involvement within these abnormal tracts.

2. Materials and methods

2.1. Subjects

Participants gave written informed consent prior to participation, and all study protocols were approved by the Institutional Review Board at the Washington University School of Medicine in St. Louis, MO. Imaging data was acquired from 18 to 30-year-old individuals, who were divided into three subject groups: 30 healthy control (CN), 24 schizophrenia (SZ) and 33 bipolar disorder (BD). Table 1 shows demographic information. Participant groups were diagnosed on the basis of a consensus between a research psychiatrist and a trained research assistant who used the Structured Clinical Interview for DSM-IV Axis I Disorder (First et al., 1996). CON subjects were required to have no lifetime history of psychotic or mood disorders. Bipolar (BD) patient participants were required to meet DSM-IV criteria for Bipolar I Disorder and were classified as psychotic BD if the patient had a psychotic event over the course of their lifetime, as assessed via the Structured Clinical Interview for the DSM (SCID). Participants were excluded if they: (a) met DSM-IV criteria for substance dependence or severe/moderate abuse during the prior 3 months; (b) had a clinically unstable or severe general medical disorder; or (c) had a history of head injury with documented neurological sequelae or loss of consciousness. Additionally, to minimize clinical heterogeneity within the BD group, only participants with a history of euphoric mania (versus mania characterized by primarily irritable mood) were included in the study.

Recent symptoms (i.e. in the last two weeks) were assessed using the Scale for the Assessment of Negative Symptoms (SANS), the Scale for the Assessment of Positive Symptoms (SAPS) (Andreasen et al., 1995), and the Young Mania Rating Scale (YMRS) (Young et al., 1978). Chronic symptoms (i.e. last several years) were assessed using the Washington Early Recognition Center Affectivity and Psychosis (WERCAP) Screen, both mania (mWERCAP) and psychosis (pWERCAP) components (Hsieh et al., 2016; Mamah et al., 2014, 2016b). Psychosocial stress severity was assessed using the WERC Stress Screen (Hsieh et al., 2016; Mamah et al., 2014).

2.2. MRI scan acquisition parameters

Scans were run using a 32-channel head coil on a customized Siemens 3T “Connectom” MRI scanner, which was previously used for collecting the Human Connectome Project – Young Adult (HCP-YA) data and housed at Washington University in St. Louis (Van Essen et al., 2012). The scanning protocol used identical parameters for individual data and housed at Washington University in St. Louis (Van Essen et al., 2012). However, the overall structure of the HCP-YA protocol was consolidated to 3 total imaging sessions (rather than 4) by only acquiring a single T1-weighted (T1w) and T2-weighted (T2w) scan (rather than two of each). The 3 sessions were typically collected over a period of two days. Briefly, T1w MPRAge and T2w SPACE images were acquired at 0.7 mm isotropic resolution. Oblique axial acquisitions alternate between right-to-left and left-to-right phase encoding directions in consecutive runs. Image reconstruction used SENSE1 multi-channel (Sotiropoulos et al., 2013). A full dMRI session included 6 runs (each approximately 9 min and 50 s), representing 3 different gradient tables, with each table acquired once with right-to-left and left-to-right phase encoding polarities, respectively. Each gradient table includes approximately 90 diffusion weighting directions plus 6 b = 0 acquisitions interspersed throughout each run. Diffusion weighting consisted of 3 shells of b = 1000, 2000, and 3000 s/mm² interspersed with an approximately equal number of acquisitions on each shell within each run. The diffusion directions were obtained using a toolbox available from INRIA that returns uniformly distributed directions in multiple q-space shells. The directions were optimized so that every subset of the first M directions was also isotropic.

2.3. Image preprocessing

Diffusion scans were concatenated and run through FSL EDDY (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/EDDY), which reduces distortions caused by eddy currents and subject movement by registering the diffusion-weighted images to the b0 image. The b0 image was registered to the T1-weighted image with an affine registration. After the distortion correction, the scans were divided to their respective polarities and averaged together. Each individual’s T1-weighted image was registered to the 1 mm-resolution MNI-152 atlas using affine registration (Jenkinson et al., 2002) in order to compare individual subjects (Collins et al., 2004). FSL’s brain extraction tool was used to remove the skull and other non-brain tissue. FSL’s “diftit” was applied to perform least-squares tensor estimations, specifically eigenvectors, eigenvalues and FA. “Bedposts GPU” (Hernandez et al., 2013) was used to apply the ball & stick model to estimate diffusion distributions.

2.4. TRACULA

TRACULA (TRActs Constrained by UnderLying Anatomy) is a powerful method (Yendiki et al., 2011) used to analyze global probabilistic tractography. This method uses a Bayesian framework for global tractography suggested by (Jbabdi et al., 2007) which determines the connection that best fits two selected endpoints using diffusion data. In addition, TRACULA also incorporates prior anatomical knowledge based on manually verified trajectories of tracts in the training set tested by (Yendiki et al., 2011). The anatomy of the tracts in the training subject set have been verified to not significantly deviate from those found in Yendiki’s clinical group (Yendiki et al., 2011). For every subject, this method reconstructs probabilistic distributions of 18 major white matter tracts in each subject. A sample participant’s estimation and identification of white matter tracts is shown in Fig. 1. More specifically, TRACULA uses the endpoints established in the training set’s tracts, expands the endpoints, and transforms them into each subject’s native space. Then, TRACULA establishes probabilistic streamlines accounting for the anatomical Freesurfer segmentations and uses control points to dictate the curvature of the tract. This method does not presume exact tract spatial location or shape, so the trajectory of the tract is only restricted with respect to the surrounding anatomical structures. This allows for individual variations across subjects while still establishing the same tracts for comparison. TRACULA trac-all package automates the segmentation steps and includes the training subject data that the manually inspected tract specifications are based on (refer to https://surfer.nmr.mgh.harvard.edu/fswiki/Tracula for more details).

We analyzed the average Fractional Anisotropy (FA) for all tracts. FA is a commonly used DTI metric that establishes the directional asymmetry of water diffusion at each voxel and total diffusion at each voxel respectively (Adler et al., 2006; Cao et al., 2014; Nortje et al., 2013; Sprooten et al., 2016). The mean FA for each subject’s tracts was calculated by TRACULA based on the probabilistic fibers that were at least 20% of the maximum probability fiber for each pathway (Yendiki et al., 2011).
2.5. TBSS

Tract-based spatial statistics (TBSS) was used to perform voxel-wise analyses of all white matter tracts, as previously described (Arnedo et al., 2015; Smith et al., 2006). FA images were calculated and projected onto the mean FA skeleton, which represents the center of white matter tracts, and thresholded at FA = 0.2.

2.6. Statistical analysis

All statistical analyses were done using SAS 9.4 (SAS Institute Inc., Cary, NC). For TRACULA studies, analysis of variance, covaried for age and sex, was used to test for diagnostic effects for each tract as well as all pairwise comparisons between diagnostic groups. Tests were Bonferroni corrected for multiple comparisons. Relationships between mean FA for each tract and clinical measures (i.e. SAPS, SANS, YMRS, mWERCAP, pWERCAP, and WERC Stress Screen) were investigated using a forward stepwise multiple linear regression analysis. We performed separate regressions to separately examine the relationship between mean FA and each behavioral variable and report the partial correlation for each significant predictor.

For voxel-wise DTI parameter analyses (i.e. TBSS), groups were compared using general linear models, covarying for age and gender. Multiple comparison corrections were applied using a permutation-based statistical approach within FSL’s Randomise (Smith et al., 2006). Randomise is FSL’s tool for nonparametric permutation inference on neuroimaging data, and produces a test statistic image and sets of P-value images (stored as 1-P for more convenient visualization, as bigger is then “better”). Brain and behavioral measures which had significant group effects or were abnormal compared to clinical norms were selected for further correlational analysis within the SZ and BD groups, controlling for age and gender. Significance was set at p < .05.

**Table 2**

FA of white matter tracts in control, bipolar disorder and schizophrenia subjects.

| Tract                             | CON   | BPD   | SCZ   | F     | p     |
|----------------------------------|-------|-------|-------|-------|-------|
| Corpus callosum                  |       |       |       |       |       |
| F. Major                         | 0.60 (0.07) | 0.59 (0.07) | 0.59 (0.08) | 0.23  | 0.79  |
| F. Minor                         | 0.54 (0.06) | 0.52 (0.08) | 0.51 (0.10) | 0.70  | 0.50  |
| Anterior thalamic radiation      |       |       |       |       |       |
| Left                             | 0.42 (0.04) | 0.41 (0.03) | 0.39 (0.06) | **3.38** | 0.039* |
| Right                            | 0.41 (0.04) | 0.41 (0.04) | 0.39 (0.06) | 1.99  | 0.14  |
| Cingular-angular bundle          |       |       |       |       |       |
| Left                             | 0.37 (0.07) | 0.40 (0.07) | 0.37 (0.08) | 1.70  | 0.19  |
| Right                            | 0.39 (0.08) | 0.41 (0.07) | 0.38 (0.08) | 0.78  | 0.46  |
| Cingulum cingulate gyrus         |       |       |       |       |       |
| Left                             | 0.59 (0.10) | 0.61 (0.08) | 0.60 (0.10) | 0.37  | 0.69  |
| Right                            | 0.54 (0.07) | 0.54 (0.06) | 0.53 (0.08) | 0.12  | 0.88  |
| Cortico-spinal tract             |       |       |       |       |       |
| Left                             | 0.54 (0.04) | 0.54 (0.04) | 0.52 (0.06) | 1.61  | 0.21  |
| Right                            | 0.52 (0.04) | 0.53 (0.05) | 0.51 (0.05) | 0.87  |       |
| Inferior longitudinal fasciculus |       |       |       |       |       |
| Left                             | 0.48 (0.04) | 0.49 (0.05) | 0.46 (0.06) | 2.58  | 0.08  |
| Right                            | 0.48 (0.04) | 0.50 (0.05) | 0.46 (0.06) | **3.64** | 0.03* |
| Sup. longitudinal fasc. - parietal|       |       |       |       |       |
| Left                             | 0.47 (0.05) | 0.48 (0.06) | 0.47 (0.06) | 0.17  | 0.84  |
| Right                            | 0.47 (0.05) | 0.47 (0.04) | 0.45 (0.06) | 1.34  | 0.27  |
| Sup. longitudinal fasc. - temp.  |       |       |       |       |       |
| Left                             | 0.49 (0.05) | 0.51 (0.04) | 0.48 (0.06) | 1.83  | 0.17  |
| Right                            | 0.47 (0.04) | 0.47 (0.05) | 0.46 (0.06) | 0.61  | 0.54  |
| Uncinate fasciculus              |       |       |       |       |       |
| Left                             | 0.37 (0.07) | 0.39 (0.07) | 0.37 (0.08) | 0.62  | 0.54  |
| Right                            | 0.38 (0.05) | 0.38 (0.07) | 0.38 (0.07) | 0.13  | 0.88  |

Bolded and asterisked values are statistically significant.
3. Results

3.1. TRACULA group analysis

Mean tract FA in each group, and post-hoc univariate tests for each tract are presented in Table 2. The most notable group differences were seen for the left anterior thalamic radiation ($p = .039$) and the right inferior longitudinal fasciculus ($p = .03$), with the lowest FA in the SZ group. These differences however did not meet statistical significance after Bonferroni correction.

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Fig. 2. $Z$-Scores of White Matter Tract Fractional Anisotropy obtained from TRACULA. Graphs depict corrected mean FA $z$-scores of each group's white matter tracts, obtained using TRACULA. $Z$-scores were corrected for gender and age, and subtracted from mean values in the control group. Figures (A) and (B) depict left and right hemispheric structures respectively, with the exception of the corpus callosum (CC) which have major and minor divisions. Plotted values represent means per group. Control = black. Schizophrenia = red. Bipolar disorder = blue. * $p < .05$. 

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Table 2: Mean tract FA in each group, and post-hoc univariate tests for each tract.
Fig. 2 depicts the z-scored (corrected) mean FA in all tracts across groups and shows a trend for SZ subjects to have lower FA compared to CON and BP participants. BP participants showed a trend towards higher tract FA compared to controls. An ANOVA comparing the 18 average tract z-scores across the three groups showed a significant effect ($F = 4.35; p = .018$). A post-hoc analysis showed significant average z-score differences between SZ and BP ($p = .01$), and trend level differences between SZ vs. CN ($p = .08$). CN vs. BP differences were not significant ($p = .3$).

### 3.2. TBSS analysis

As shown in Fig. 3A, compared to control subjects, significantly decreased FA was observed in SZ patients largely in the left cerebral hemisphere, unilaterally distributed over the anterior thalamic radiation, posterior thalamic radiation, corticospinal tract and superior longitudinal fasciculi (FDR correction $p < .05$). Bilaterally decreased FA in the cerebellum was also observed in SZ patients compared to controls.

Compared to controls, FA in bipolar disorder patients were not statistically different. Group FA differences between BP and SZ patients were bilateral and widespread, with decreased FA in SZ patients (Fig. 3B).

### 3.3. Clinical relationships: TRACULA

In order to assess the relationship between clinical variables and FA of TRACULA tracts, across BP and SZ participants, we performed separate stepwise linear regressions for chronic mania (mWERCAP), chronic psychosis (pWERCAP), stress (WERC Stress Screen), recent manic symptoms (YMRS), recent positive symptoms (SAPS), recent disorganization (SANS), and negative symptoms (SANS), respectively. Among all white matter tracts, the regression for chronic mania identified only the anterior thalamic radiation (ATR) FA as predicting chronic mania severity, accounting for 19.9% of the variance ($B = 38.5; F = 13.7; p = .0005$). A separate multiple regression showed only the cingulum-angular bundle (CAB) FA significantly predicted recent mania, accounting for 9.3% of the variance ($B = 6.7; F = 5.66; p = .02$). No tract predicted any other behavioral symptom across BP and SZ participants.

Multiple regression analyses were also conducted separately in BP participants (mWERCAP, YMRS) and SZ participants (pWERCAP, SAPS, SANS). In BP, the superior longitudinal fasciculus-temporal (SLF-T) FA predicted chronic mania with 12% of the variance ($B = −23.9; F = 4.2; p = .049$), and the CAB FA predicted recent mania with 12.9% of the variance ($B = 9.7; F = 4.6; p = .04$). In SZ, the only CAB FA predicted chronic psychosis with 23.5% of the variance ($B = 29.5; F = 6.8; p = .016$).

### 3.4. Clinical relationships: TBSS

We calculated the relationship between each behavioral measure with FA values across both BP and SZ participants.

Among all behavioral measures, only chronic mania (mWERCAP) correlated with FA, specifically within the left anterior corpus callosum and left anterior thalamic radiation (Fig. 4). Increased FA was related to increased chronic mania severity.

### 3.5. Medication relationships

Considering the highly heterogenous medications (and medication combinations) used across our patient populations, completely controlling for medications was not possible. We however investigated the effects of more commonly used medications that have been associated with structural brain changes in select patient groups: 1) typical antipsychotics vs. atypical antipsychotics vs. neither in SZ; 2) lithium vs. no lithium in BP; 3) atypical vs. no atypical in BP; and 4) SSRI/SNRI in BP. Fig. 5 shows results of individual comparisons and mean tract FA values across groups. Notably, the SSRI/SNRI group trended towards lower mean FA than the non-SSRI/SNRI group, with significant findings observed in the ATR ($F = 6.8; p = .014$); SLF-P ($F = 4.3; p = .046$); SLF-T ($F = 4.7; p = .039$). Group differences did not remain after correcting for multiple comparisons.

### 4. Discussion

We investigated white matter integrity in the brain of patients with schizophrenia and bipolar disorder, scanned with an optimized
The atrophy of the cingulum-angular bundle (CAB) is significant in schizophrenia patients, although with some variability in the affected tracts across studies. Decreased structural connectivity of the cingulum, primarily in its anterior part, has been associated with bipolar disorder (Ji et al., 2017). Decreased FA in the temporal tract of the superior longitudinal fasciculus (SLF-T) also predicted chronic affective symptoms in bipolar patients suggesting that impaired integrity of this tract may be related to the neurobiology of the illness. The SLF is a widespread white matter tract in this disorder and specific medication influences.

A multiple regression identified the cingulum-angular bundle (CAB) FA as predicting recent mania severity in bipolar patients alone, as well as across both patient groups. Interestingly, the CAB FA also predicted chronic psychotic symptoms in patients. In both cases, increased FA in the CAB correlated with increased symptoms. The CAB represents the posterior part of the cingulum, which extends between the posterior cingulate gyrus and the parahippocampal and uncus of the temporal lobe (Jones et al., 2013). Fibers within this tract have been associated with cognitive functions, conveying memory information from the hippocampus and integrating it with other parts of the brain (Ezzati et al., 2016). Decreased structural connectivity of the cingulum, primarily in its anterior part, has been associated with bipolar disorder (Ji et al., 2017; Wang et al., 2008), and with mania (Martino et al., 2016). Our study is the first to our knowledge reporting a positive correlation between cingulum integrity and manic severity, and psychosis in schizophrenia. While speculative, this may reflect enhanced fiber coherence related to prolonged hippocampal and/or amygdala hyperactivity, which has been associated with bipolar disorder (Heckers and Konradi, 2015; Tregellas et al., 2014). Decreased FA in the temporal tract of the superior longitudinal fasciculus (SLF-T) also predicted chronic affective symptoms in bipolar patients suggesting that impaired integrity of this tract may be related to the neurobiology of the illness. The SLF is a widespread white matter tract in this disorder and specific medication influences.
major association tract in the brain that connects frontal, occipital, parietal and temporal lobes of the brain, and was the only white matter tract shown to have decreased FA in both existing TRACULA studies of bipolar disorder (Ji et al., 2017; Sprooten et al., 2016). Specifically, the impaired SLF-T integrity was also related to psychotic symptoms in one of these studies (Ji et al., 2017), suggesting this tract may be a marker of illness severity and not of a specific symptom.

The implications of FA abnormalities in disease has been well described. FA measures the asymmetry in the direction of diffusion of water. To the extent that white matter fiber tracts are structurally intact and healthy, the diffusion of water will be dominated by a coherent direction down the path of the fiber tract (Beaulieu, 2002). Axonal degradation, demyelination, and neurodegenerative effects however would be expected to produce a disruption in the coherence of water diffusion and consequently a decrease in the FA. Such findings appear to be largely related to underlying neurobiology of schizophrenia, considering low FA is often seen in medication naïve patients (Cheung et al., 2008, 2011; Filippi et al., 2014; Gasparotti et al., 2009; Guo et al., 2012; Perez-Iglesias et al., 2016; Sun et al., 2015) and unaffected siblings or high-risk individuals (Epstein et al., 2014; Peters et al., 2009; von Hohenberg et al., 2014; Zhou et al., 2017). Nevertheless, the effects of confounding factors, including medications and substance use which are prevalent in these patients, may influence the pattern of white matter architecture. Differences in scan acquisition, motion correction, and analysis methods could also contribute to variability across studies. Beyond technical issues and external factors, the biological heterogeneity of schizophrenia likely manifest in differential patterns of brain connectivity, sometimes with differential symptomatology (Arnedo et al., 2015; Sun et al., 2015). For example, by unsupervised biclustering of TBSS-derived white matter data, Arnedo et al. (2015) identified four general patterns of low FA among schizophrenia subjects, predominantly involving the genu of the corpus callosum; fornix and external capsule; splenium of the corpus callosum; and anterior limb of the internal capsule respectively. Along these lines, Sun et al. (2015) applied hierarchical clustering of 36 features from fiber tracts in schizophrenia patients and found two patterns of abnormalities: one showing widespread white matter abnormalities and another with primarily superior longitudinal fasciculi abnormalities. Thus, it is likely that any given cohort of schizophrenia patients comprises of a highly heterogenous combination of participants, making broad conclusions about specific schizophrenia abnormalities impractical.

There are some limitations in our ability to interpret and extend upon the current findings. Due to the cross-sectional nature of the current study, we cannot address questions of causality in the relationship of observed white matter fractional anisotropy abnormalities. Therefore, this study is unable to address the time course of FA changes with bipolar disorder and cannot specifically rule out the effects of various forms of treatment over the course of the disease. As alluded to above, we also cannot account for all of the potential medication effects on white matter integrity in the current sample, as our study was not well powered to investigate medication effects. For example, many of our patients were taking various mood stabilizers, which could potentially have influenced structural connectivity. Specifically, the neuroprotective effects of mood stabilizing pharmaceuticals may counteract impaired connectivity results differentially throughout the brain in various samples. Lithium, for example, has been associated with the regulation of cell death as well as up-regulation of proteins responsible for axonal myelination, a vital property of

![Fig. 5. Medication Effects on TRACULA-Derived White Matter Tracts. Figures depict mean group FA z-scores, corrected for gender and age. Left and right tracks have been combined. (A) schizophrenia patients on either typical antipsychotics, atypical antipsychotics or neither; (B) bipolar disorder patients either on selective serotonin reuptake inhibitor (SSRI)/serotonin norepinephrine reuptake inhibitor (SNRI) or not; (C); bipolar disorder patients either on lithium or not. * p < .05.]
cell growth that would directly affect measures of white matter integrity (Chen et al., 1999; Manji et al., 2000; Nonaka et al., 1998). Antipsychotic medications, used in the treatment of schizophrenia and often bipolar disorder, have been associated with decreased white matter FA (Szeszko et al., 2014; Wang et al., 2013) and volume (Girgis et al., 2006; Ho et al., 2011), possibly from alterations in glial cell composition or number (Szeszko et al., 2014). Indirectly, white matter connectivity could also be affected indirectly, though medication effects on cortical gray matter size (Lieberman et al., 2005). Finally, step-wise linear regression may perform poorly with the number of predictors included in cases of collinearity and could give biased coefficients that need shrinkage (Tibshirani, 1996).

In summary, we investigated white matter integrity in schizophrenia and bipolar disorder using two complementary approaches. Both approaches showed reduced white matter FA in schizophrenia, with a trend towards increased FA in bipolar disorder, compared to controls. Specifically, we found decreased FA in multiple, primarily left sided, white matter regions in schizophrenia using TBSS, a voxel-based approach. A tract-based methodology, TRACULA, showed a trend towards lower FA in multiple tracts in schizophrenia, and a trend towards higher FA in some tracts in bipolar disorder. FA in the cingulum-angular bundle predicted chronic mania, and in the superior longitudinal fasciculus predicted recent mania. These studies build on existing studies of psychiatric disorders and show decreased white matter integrity in schizophrenia. Medications, drug use, age and biological heterogeneity are likely confounds, and may have contributed to absence of findings in bipolar patients. Future studies using larger groups are needed, and longitudinal investigations of effects of external factors on structural brain connectivity.

Funding

This work was funded by NIMH grant R01 MH104414. Additionally, Dr. Mamah has received funding from Taylor Foundation Institute, Dept. Psychiatry, Washington University and the Center for Brain Research on Mood Disorders, Dept. Psychiatry, Psychiatry University. Research reported in this publication was also supported by the Eunice Kennedy Shriver National Institute of Child Health & Human Development of the National Institutes of Health under Award Number U54 HD087011 to the Intellectual and Developmental Disabilities Research Center at Washington University.

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