

Frontal White Matter Integrity in Borderline Personality Disorder With Self-Injurious Behavior

Jon E. Grant, J.D., M.D.
 Stephen Correia, Ph.D.
 Thea Brennan-Krohn, B.A.
 Paul F. Malloy, Ph.D.
 David H. Laidlaw, Ph.D.
 S. Charles Schulz, M.D.

Self-injurious behavior in borderline personality disorder is a frequent cause of morbidity and mortality, but neurobiological studies examining this behavior are few. Nine women with borderline personality disorder self-injurious behavior and seven comparison subjects underwent diffusion tensor imaging (DTI). Trace and fractional anisotropy (FA) were calculated for frontal and posterior regions. Borderline personality disorder-self-injurious behavior subjects also underwent a battery of neuropsychological tests that emphasized executive functions. They had significantly higher trace and lower FA in inferior frontal but not posterior regions. Correlational analyses between DTI and cognitive variables showed a pattern of results that was contrary to expectations with posterior white matter integrity correlating with isolated measures of executive function and anterior white matter integrity correlating with a component of verbal memory test performance. Women with borderline personality disorder-self-injurious behavior exhibit decreased white matter microstructural integrity in inferior frontal brain regions that may include components of orbito-frontal circuitry.

(The Journal of Neuropsychiatry and Clinical Neurosciences 2007; 19:383–390)

Approximately 75% of individuals diagnosed with borderline personality disorder engage in impulsive self-injurious behavior,¹ and this behavior is a frequent cause of emergency room visits, hospitalizations, and premature morbidity and mortality. The impulsive aggression seen in this subgroup of individuals with borderline personality disorder may be associated with frontal lobe dysfunction and may arise from impaired connectivity between the frontal lobe and other regions.^{2–4}

Imaging studies of individuals with borderline personality disorder are few.⁵ Early studies using computed tomographic imaging reported no gross abnormalities, no difference in ventricle-brain ratio, and no evidence of frontal lobe atrophy in borderline personality disorder patients compared with healthy subjects.⁶ A recent magnetic resonance image (MRI) volumetry study found evidence of significantly reduced volumes of left orbito-frontal and right anterior cingulate cortex in borderline personality disorder subjects compared with healthy

Received July 17, 2006; revised October 30, 2006; accepted November 20, 2006. Drs. Grant and Schulz are affiliated with the University of Minnesota, Minneapolis, Minnesota. Dr. Correia is affiliated with the Providence Veterans Affairs Medical Center, Brown Medical School, Providence, Rhode Island. Ms. Brennan-Krohn and Dr. Malloy are affiliated with Butler Hospital, Brown Medical School, Providence, Rhode Island. Dr. Laidlaw is affiliated with Brown University, Providence, Rhode Island. Address correspondence to Dr. Grant, Department of Psychiatry, University of Minnesota Medical Center, 2450 Riverside Avenue, Minneapolis, MN 55454; grant045@umn.edu (e-mail).

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subjects.⁷ Positron emission tomography studies of borderline personality disorder have resulted in conflicting results of both hypo- and hyper- frontal metabolism.^{8–10} A functional MRI study showed greater activation in the amygdala, and medial and inferolateral prefrontal cortex in borderline personality disorder subjects compared with healthy subjects, reflecting more intense emotional responses to stressors.¹¹

Diffusion tensor imaging (DTI) is an MRI technique that measures the magnitude and direction of water diffusion in brain tissue. DTI data can be visualized in a variety of ways, including two-dimensional maps of the scalar parameters: a) trace, a measure of the magnitude of water diffusion in each image voxel; and b) fractional anisotropy (FA), a measure of the extent to which water diffusion in each voxel is directionally restricted. Typically, in regions of compromised white matter integrity, trace values are higher and FA values lower than in normal white matter, presumably owing to axonal degeneration.¹²

Prior DTI studies have demonstrated an association between impulsivity and the reduction of microstructural integrity of frontal white matter systems. For example, Hoptman et al.^{4,13} found that lower FA (i.e., axonal disorganization) in the right inferior frontal white matter was associated with greater impulsivity in schizophrenia. Additionally, our own work using DTI has shown decreases in the structural integrity of frontal but not posterior white matter in patients with kleptomania.¹⁴

Because frontal brain circuits, particularly the orbitofrontal circuit, are important in behavioral regulation,¹⁵ we hypothesized that individuals with borderline personality disorder who engage in self-injurious behavior would show compromised white matter integrity (i.e., increased trace and decreased FA) in inferior frontal regions, but not in posterior regions, compared with a healthy comparison group using DTI. We also hypothesized that anterior white matter integrity measured by trace and FA would be correlated with executive functions but not with cognitive functions, such as naming and basic visuospatial perceptual ability, which are thought to be more dependent on posterior cortical regions.

METHOD

We recruited 10 women with borderline personality disorder self-injurious behavior (mean age = 34.1 [SD =

10.8]; range = 18 to 51; all right-handed) from an outpatient clinic. The diagnosis was confirmed by the Structured Clinical Interview for DSM-IV Personality Disorders (SCID-II).¹⁶ Because borderline personality disorder may have multiple domains with distinct neurobiological underpinnings,¹⁷ we restricted our study to only on those borderline personality disorder subjects who reported an inability to control their impulses to self-injure. Therefore, inclusion criteria were: 1) borderline personality disorder as the primary psychiatric disorder; 2) self-injurious behavior (defined as impulsive acts of self-mutilation, such as cutting, head banging, or burning) at least once a week; and 3) urges to self-injure at least one time per week. Subjects rated intensity of urges to self-injure using a 10-point Likert scale (0 = no urges; 10 = incapacitating urges). We recruited 10 healthy, non-psychiatric female subjects (mean age = 32.8 [SD = 9.5]; range = 21 to 49; all right-handed) matched to the borderline personality disorder-self-injurious behavior group on key demographic variables from the community.

Exclusion criteria for all subjects included: 1) current or lifetime history of bipolar I or psychotic disorder based on the Structured Clinical Interview for DSM-IV (SCID);¹⁸ 2) a lifetime history of attention deficit hyperactivity disorder based on clinical interview; 3) a lifetime history of a DSM-IV impulse control disorder not elsewhere classified based on SCID-compatible modules; 4) a history of head injury or neurological disorder; and 5) a positive urine pregnancy test.

Subjects taking psychotropic medications were allowed to participate if the dose had been stable for at least 6 months prior to study entry and had not resulted in any subjective improvement in self-injurious urges or behavior.

Butler Hospital's Institutional Review Board approved the study. After complete description of the study, subjects provided written informed consent.

Assessments

We evaluated subjects at entry into the study by the Structured Clinical Interview for DSM-IV (SCID)¹⁸ and SCID-compatible modules for impulse control disorders.¹⁹ Self-injurious behavior was assessed with a semi-structured phenomenological questionnaire evaluating types of self-injurious behaviors, frequency of behaviors, and related emotions before, during, and after self-injury. In addition, depressive symptoms were assessed using the Hamilton Depression Rating Scale (HAM-D),

a valid and reliable 17-item, clinician-administered rating scale evaluating the severity of depressive symptoms.²⁰

Neuropsychological Battery

Borderline personality disorder-self-injurious behavior subjects underwent a battery of neuropsychological tests that emphasized executive functions. Two of the 10 subjects refused to undergo neuropsychological testing due to time constraints. The battery was administered by a clinical neuropsychologist. The tester was aware of the subjects' diagnoses. Testing duration was approximately 1.5 to 2 hours. Control subjects were not administered the battery as normative means based on gender and age have been published.

Procedures

MRI scans were obtained on a 1.5T Siemens Symphony scanner using a volume head coil. A standard localizer was obtained followed by a 3D T1 MPRAGE (one acquisition, sagittal) as follows: 0.85 mm slices, no gap, 176 slices, 256×256 matrix, 21.7×21.7 cm FOV, TR=1900, TE=4.31 msec, TI=1100, NEX=1, and flip angle=15; acquisition time=8.08 minutes. Coregistered sagittal double spin-echo, echo-planar diffusion-weighted images were collected based on Siemens' MDDW protocol as follows: three acquisitions with offset in slice direction by 0.0 mm, 1.7 mm, and 3.4 mm; 5 mm thick slices; 0.1 mm interslice spacing; 30 slices per acquisition; 128×128 matrix, 21.7 cm×21.7 cm FOV (interleaving during post-processing provides true 1.7 mm³ resolution images), TR=7200, TE=156. Bipolar diffusion gradients were applied in 12 noncollinear diffusion directions with 2 b magnitudes: 0, 1000 mm/s², NEX=3, no partial echoes. A double-echo sequence was used that effectively cancels eddy current effects.²¹ The entire brain was imaged. Time per acquisition=4:48 minutes. We used a vacu-pillow and head cushions to minimize subject movement during scanning.

All three offset diffusion scans were up-sampled to 0.85 mm³ isotropic voxels for analysis. Scalar maps of trace and FA were produced using custom software.²² An additional T2-weighted image (I0) without diffusion encoding (b=0) inherently coregistered with the trace and FA images was also produced.

Image Analysis

DTI data on one borderline personality disorder-self-injurious behavior subject and three comparison subjects

could not be analyzed due to motion artifact. The final sample included nine borderline personality disorder-self-injurious behavior patients and seven comparison subjects. The mean age was 33.7 (SD=11.3) years (range: 18 to 51 years) in the borderline personality disorder-self-injurious behavior group and 31.1 (SD=10.6) years (range: 21 to 49 years) in the comparison subjects. This difference in age was not statistically significant ($t = -0.455$, $p = 0.656$); skewness and kurtosis were within expectations for the age distribution for either group.

Experienced raters (S.C. and T.B.K.), blind to group assignment, analyzed images using Analyze AVW software (v. 5.0 & 6.0).²³ The MPRAGE images were manually corrected for head rotation and resliced along the AC-PC line. The transform matrix was applied to the DTI FA map volumes with manual adjustment and then this adjusted matrix was applied without further adjustment to the remaining DTI maps (trace and b=0). Four standard sized square (5 mm×5 mm voxel) regions of interest (ROIs) were placed bilaterally in anterior and posterior white matter on each of four axial slices based on a previously published method³ for a total of 16 regions per subject. The most inferior slice was identified in sagittal view and was located at the inferior border of the rostrum of the corpus callosum. The remaining three slices were those falling three, six, and nine slices superior to the first. A prespecified coordinate-based algorithm was designed to guide ROI placement such that anterior ROIs would be placed anterior and slightly lateral to the anterior horns of the lateral ventricles on the three superior slices, and anterior and medial to the Sylvian fissure on the most inferior slice, and posterior ROIs would be placed lateral to the posterior horns of the lateral ventricles. Adjustments in final ROI placement were made to accommodate individual differences in brain anatomy. All ROIs were placed on the b=0 image without reference to the trace and FA images and transferred without further adjustment to the inherently coregistered trace and FA images for measurement.

The 16 ROIs were placed by each rater on each of the 16 brain volumes (nine patients, seven comparison subjects), for a total of 16 measurements per ROI per rater (a total of 256 ROIs placed per rater). Across the 16 regions, interrater reliability, measured as the intraclass correlation coefficient (ICC) between two raters, ranged from 0.71 to 0.98 for trace with only two regions falling below 0.80; and ranged from 0.78 to 0.98 for FA with only one region falling below 0.80. To optimize mea-

surement reliability, we omitted those individual FA and trace measurements that differed more than 15% between the raters. After removal of these individual measurements, ICC improved to 0.84 or greater for 15 of the 16 trace ROIs and was 0.76 for the remaining region; ICC improved to 0.86 or greater for the 16 FA ROIs. To further improve reliability, we used the mean of the two raters' values for trace and FA for each of the 16 regions in the analysis. For the group analysis, the 16 ROIs (i.e., right and left anterior and posterior regions on each of four slices) were summed across slices so that each subject had four trace and four FA measurements (i.e., right and left anterior and right and left posterior).

Cognitive Testing

Borderline personality disorder-self-injurious behavior subjects underwent a battery of neuropsychological tests that emphasized executive functions: Behavioral Dyscontrol Scale (BDS)²⁴ for executive functioning; Trail-Making Test, Parts A and B (TMT-A and -B)²⁵ for executive functioning, sequencing; psychomotor processing speed, cognitive set switching; Controlled Oral Word Association Test (COWAT)²⁶ for executive functioning, lexical fluency; Stroop Color-Word Test (Stroop)²⁷ for response inhibition; Wisconsin Card Sorting Test (WCST)²⁸ for executive functioning, mental flexibility; Boston Naming Test (BNT)²⁹ for language, confrontational naming; Judgment of Line Orientation (JLO)³⁰ for visuospatial perception; Hopkins Verbal Learning Test-Revised (HVLT-R)³¹ for memory, verbal list learning and recall.

Data Analysis

Student's *t* tests and two-group one-way ANOVA were used to examine group differences in background variables. Group differences in trace and FA were analyzed in separate (2 × 2) repeated measures (one for trace and one for FA) ANOVA with group (borderline personality disorder versus comparison) as the between-groups factor and region (anterior-posterior) as the within-groups factor. Follow-up analysis of simple effects was performed using one-way ANOVA. We did not use age as a covariate in our repeated measures analyses given that the means and frequency distributions of age in the two groups were highly comparable. There were no right versus left hemisphere differences in trace or FA in either group, with the exception of significantly greater posterior FA on the left in the borderline personality disorder-self-injurious behavior group only ($F [1, 70] =$

4.21, $p = 0.04$). However, since we did not pose a hemispheric lateralization hypothesis, we summed anterior and posterior ROIs across hemispheres. The relationship between anterior and posterior trace and FA values and performance on cognitive tests was assessed in the borderline personality disorder-self-injurious behavior group using Pearson bivariate correlations. We used age- and education-corrected *T* scores for this analysis and since our hypothesis was directional, we used a one-tailed test. We used an alpha level of $p < 0.05$; we did not adjust the alpha level to reflect multiple comparisons because this is the first study of this topic and is therefore exploratory.

RESULTS

The demographics of the borderline personality disorder-self-injurious behavior and comparison groups are presented in Table 1. The borderline personality disorder-self-injurious behavior group was significantly more likely to have graduated from college and have a history of major depressive disorder, and higher scores on the HAM-D. The groups did not differ significantly on other demographic variables.

Mean self-injurious behavior onset was 16.7 (SD = 4.1) years (range = 9 to 23). Mean self-injurious behavior urge intensity was 7.40 (SD = 1.58), and mean self-injurious behavior frequency was 3.40 (SD = 1.17) times per week at the time of assessment. Eight subjects had been psychiatrically hospitalized due to self-injurious behavior, and three had required medical care (e.g., stitches, antibiotics). All subjects cut themselves, and six reported additional self-injurious behavior (burning [N = 3]; head banging [N = 2], beating self [N = 1]). Lifetime comorbidity data are presented (Table 1). Although only three of the borderline personality disorder-self-injurious behavior subjects met criteria for lifetime PTSD, all nine had histories of sexual abuse compared to none in the comparison group.

The nine borderline personality disorder-self-injurious behavior women had extensive treatment histories. All nine women had experienced at least 1 year of group dialectical behavioral therapy (DBT) plus therapy with an individual DBT counselor. Seven of the nine women had undergone 2 or more consecutive years of DBT groups. Mean onset of treatment was at 21.7 (SD = 8.4) years (range = 13 to 25). All of the women were currently taking at least one psychotropic medication (mean num-

ber of current psychotropic medications per subject was 4.0 (SD=1.3 [range=2 to 6]). Of the medications currently prescribed for the nine women, seven were taking at least one atypical antipsychotic, six were taking at least one antidepressant and at least one hypnotic, five were taking at least one mood stabilizer, and one was taking a stimulant.

DTI Results

Repeated measures ANOVA with diagnosis as the between-subjects factor and region (anterior-posterior) as the within-subjects factor revealed significant main effects of group ($F [1, 14] = 7.81, p = 0.014$) and region ($F [1, 14] = 14.19, p = 0.002$). There was a significant group-by-region interaction effect ($F [1, 14] = 7.30, p = 0.017$). Follow-up analysis of simple effects revealed that this interaction was driven by significantly higher anterior trace in the borderline personality disorder-self-

injurious behavior group compared with the comparison group ($F [1, 14] = 10.45, p = 0.006$); the groups did not differ significantly in posterior trace ($F [1, 14] = 0.205, p = 0.658$) (Table 2).

A second repeated measures ANOVA revealed nonsignificant trends for the main effects of group ($F [1, 14] = 4.29, p = 0.057$) region ($F [1, 14] = 3.48, p = 0.083$). There was a significant group-by-region interaction effect ($F [1, 14] = 9.92, p = 0.007$). Follow-up analysis of simple effects revealed that this interaction was driven by significantly lower anterior FA in the borderline personality disorder-self-injurious behavior group compared with the comparison group ($F [1, 14] = 11.39, p = 0.005$); the groups did not differ significantly in posterior FA ($F [1, 14] = 0.167, p = 0.689$) (Table 2).

Neither self-injurious behavior frequency nor self-injurious behavior urge intensity was significantly correlated with either anterior trace ($|r| = -0.193, p = 0.620$);

TABLE 1. Demographics and Clinical Characteristics of Women With Borderline Personality Disorder and Self-Injurious Behavior (BPD-SIB) Compared With Healthy Subjects

	BPD-SIB (N=9)	Healthy Subjects (N=7)	Statistic	df	p value
Age			0.286t	14	0.778
Mean (SD),	34.1 (10.8)	32.8 (9.5)			
Range in years	18 to 51	21 to 49			
Ethnicity, N (%)					
Not Hispanic/Latino	9 (100)	7 (100)			
Marital status, N (%)			4.216c	2	0.121
Single	7 (77.8)	3 (42.9)			
Married	1 (11.1)	4 (57.1)			
Widow/separated/divorced	1 (11.1)	0 (0)			
Education, N (%)			-2.474z	n/a	0.013
High school grad or less	0 (0)	2 (28.6)			
Part college	1 (11.1)	3 (42.9)			
College grad	7 (77.8)	2 (28.6)			
Postcollege education	1 (11.1)	0 (0)			
Unemployed, N (%)	2 (22.2)	1 (14.3)	f	n/a	1.0
HAM-D total score, mean (SD)¹	11.9 (3.9)	0.4 (0.8)	-8.690t	14	<0.001
Comorbid lifetime disorders, N (%)²					
Major depressive disorder	8 (88.9)	2 (28.6)	f	n/a	0.035
Posttraumatic stress disorder	3 (33.3)	1 (14.3)	f	n/a	0.585
Any other anxiety disorder	1 (11.1)	1 (14.3)	f	n/a	1.0
Alcohol abuse/dependence	2 (22.2)	0 (0)	f	n/a	0.475
Drug abuse/dependence	2 (22.2)	0 (0)	f	n/a	0.475
Nicotine dependence	2 (22.2)	0 (0)	f	n/a	0.475
Any eating disorder	3 (33.3)	1 (14.3)	f	n/a	0.585
Any somatoform disorder	0 (0)	0 (0)	f	n/a	1.0
ADHD	2 (22.2)	1 (14.3)	f	n/a	0.213
Any other personality disorder	3 (33.3)	0 (0)			

For Statistic column:

t = t-test

c = Chi-square

z = Z for Mann-Whitney test

f = Fisher's exact test (two-sided)

Ethnicity and "any somatoform disorder" are constants and no statistical test was run.

¹ HAM-D = 17-item Hamilton Depression Scale

² Lifetime disorders based on SCID assessment

[$r = -0.098$, $p = 0.802$], respectively) or anterior FA ([$r = 0.206$, $p = 0.594$]; [$r = 0.219$, $p = 0.571$], respectively).

Cognitive Results

Cognitive data were available for only eight of the nine borderline personality disorder-self-injurious behavior subjects with analyzable imaging data. Briefly, the means for all scores derived from the cognitive tests were within normal limits, with the exception of a number of categories completed on the WCST (mean = 4.5 [SD = 2.14]). For the correlation analysis, we chose only cognitive variables for which the range of the scores in the sample extended into the impaired range (i.e., T score ≥ 1 SD below the mean). Posterior trace was significantly correlated ($p < 0.05$), with perseverative ($r = 0.646$) and nonperseverative ($r = 0.627$) responses on the WCST. Anterior trace and FA were significantly correlated ($r = 0.755$ and -0.753 , respectively, $p < 0.05$), with true positive responses on the HVLT-R recognition recall trial.

DISCUSSION

These DTI results appear to support our hypothesis that patients with borderline personality disorder-self-injurious behavior exhibit compromised frontal white matter systems. These findings of compromised white matter microstructure in inferior frontal regions are consistent with results reported in other impulsive behaviors and with the hypothesis that impaired inferior frontal brain circuits underlie impulsive aggressive behaviors.³ Compromised frontal white matter microstructure in patients with borderline personality disorder-self-injurious behavior may reflect an inability to balance the desire for immediate gratification from cutting with the recognition of the long-term consequences, an activity that may involve prefrontal cortical function.³²

The exploratory correlational analyses between DTI and cognitive variables in the borderline personality disorder-self-injurious behavior group yielded mixed pattern results vis-à-vis our executive cognitive hypothesis. In fact, higher posterior, not anterior, trace was associated with higher numbers of errors on the WCST, whereas higher anterior trace and lower FA was associated with better performance (i.e., higher numbers of true-positive responses) on a verbal recognition recall task. This latter finding is clearly the opposite of what was expected. These findings are inconsistent with other studies assessing neurocognitive functioning in borderline personality disorder.^{33,34} Previous studies of neurocognitive functioning in borderline personality disorder have produced mixed results with some,^{35,36} but not others^{37,38} reporting distinct impairments relative to comparison subjects. Whether inconsistent results from previous studies are due to the possible heterogeneity of borderline personality disorder is unclear. Our sample was fairly homogenous, consisting only of female borderline personality disorder patients with urges to self-injure who were treatment-resistant. Although we had no cognitive data on the comparison subjects with which to compare, no robust pattern of neuropsychological impairment emerged when compared to test norms. In fact, as a group, the borderline personality disorder-self-injurious behavior participants performed within normal limits on virtually all tests thereby limiting the ability to identify significant associations. An alternative explanation is that executive cognitive functions and impulse control are subserved by distinct frontal regions—dorsolateral and orbital, respectively. Previous reports have demonstrated that cognitive deficit and disinhibition are dissociable in patients with focal frontal lesions.^{39,40} Our analytical approach did not examine whether a similar dissociation exists in patients with borderline personality disorder-self-injurious be-

TABLE 2. White Matter Fractional Anisotropy and Trace in Women with Borderline Personality Disorder and Self-Injurious Behavior (BPD-SIB) Compared to Healthy Controls

Region and DTI parameter	BPD-SIB Subjects (SD) (N = 9)	Healthy Subjects (SD) (N = 7)	F (1,14)	p value
Anterior				
Fractional Anisotropy	0.4530 (0.0732)	0.5818 (0.0789)	11.39	0.005
Trace ($\text{mm}^2\text{s}^{-1} \times 10^{-3}$)	0.7478 (0.0921)	0.5840 (0.1109)	10.45	0.006
Posterior				
Fractional Anisotropy	0.4821 (0.0657)	0.4684 (0.0667)	0.17	0.689
Trace ($\text{mm}^2\text{s}^{-1} \times 10^{-3}$)	0.7830 (0.0813)	0.7975 (0.0265)	0.21	0.658

Trace and fractional anisotropy are presented as means (SD) by group and region (simple mean comparisons)
DTI = diffusion tensor imaging

havior. Lastly, the very small size of our sample relative to the fairly large number of neuropsychological test scores raises questions about the reliability of these cognitive/imaging correlations.

The DTI findings of compromised inferior frontal microstructure may explain why certain therapies and medications have historically proven useful for borderline personality disorder behaviors. Psychotherapies that improve self-regulation⁴¹ may counterbalance possible frontal deficits. Similarly, pharmacotherapies that increase inhibition, possibly through action in the prefrontal cortex,⁴² may reduce self-injurious behavior in borderline personality disorder.

This is a preliminary analysis, and the findings should be interpreted cautiously. First, although our acquisition protocols control for eddy current and susceptibility artifacts, such effects may have affected our measurements, particularly in anterior regions where such artifacts tend to be greater. However, it is unlikely that such effects would have interacted systematically with subject group to produce a bias favoring our hypothesis. Second, we reduced the likelihood that motion artifact had a strong impact on the results by removing from the analysis cases with excessive motion. It is unlikely that subtle motion artifact (e.g., from physiological effects) in the retained cases would have produced systematic group bias in ROI measurements. Third, the sample size was small, and therefore replication in a larger sample is warranted. Fourth, the sample was limited to borderline personality disorder-self-injurious behavior subjects who were still engaging in self-injury at a late age despite an extensive treatment history. Therefore, these subjects appear to represent a largely treatment-resistant subset of borderline personality disorder subjects. These findings, therefore, may not generalize to all borderline personality disorder-self-injurious behavior subjects. Fifth, no comparison group of borderline personality disorder subjects without self-injurious behavior were

examined. Only by using such a comparison group could we appreciate validity to the FA analysis, that is, whether disturbed frontal white matter is related to the borderline personality disorder diagnosis or the self-injurious behavior. Finally, the presence of co-occurring lifetime disorders may have contributed to the observed between-group differences in the frontal cortex. Without a comparison to another psychiatric sample with impulsivity, we cannot comment on how specific these findings are to this subject population. We did not control for age in our main analysis because the age distributions in the groups were highly similar. Moreover, there was no overall effect of age by region when age was added as a covariate in our main analysis ($p = 0.560$; data not shown). These results lessen the likelihood that age was an important confounding factor in our results, but it cannot be ruled out in this small sample.

Despite these limitations, including its restriction in sample size, this study demonstrated significant decreased white matter microstructural integrity in inferior frontal brain regions of women with borderline personality disorder-self-injurious behavior. To the extent that we sampled components of orbitofrontal white matter circuits, our results provide preliminary support for the hypothesis that orbitofrontal abnormalities may underlie some of the behavioral dysregulation in borderline personality disorder patients with self-injurious behavior. The study of a larger number of subjects with more in-depth assessments of impulsivity are the logical next steps for this research. It remains to be determined whether successful response to treatment in borderline personality disorder-self-injurious behavior individuals is dependent upon white matter integrity.

This study was supported by a Young Investigator Award from the National Alliance for Research on Schizophrenia and Depression (NARSAD) to Dr. Grant. The study was performed at Butler Hospital, Brown Medical School, Providence, RI.

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