Right Arcuate Fasciculus Abnormality in Chronic Fatigue Syndrome

Purpose: To identify whether patients with chronic fatigue syndrome (CFS) have differences in gross brain structure, microscopic structure, or brain perfusion that may explain their symptoms.

Materials and Methods: Fifteen patients with CFS were identified by means of retrospective review with an institutional review board–approved waiver of consent and waiver of authorization. Fourteen age- and sex-matched control subjects provided informed consent in accordance with the institutional review board and HIPAA. All subjects underwent 3.0-T volumetric T1-weighted magnetic resonance (MR) imaging, with two diffusion-tensor imaging (DTI) acquisitions and arterial spin labeling (ASL). Open source software was used to segment supratentorial gray and white matter to compare gray and white matter volumes and cortical thickness. DTI data were processed with automated fiber quantification, which was used to compare piecewise fractional anisotropy (FA) along 20 tracks. For the volumetric analysis, a regression was performed to account for differences in age, handedness, and total intracranial volume, and for the DTI, FA was compared piecewise along tracks by using an unpaired t test. The open source software segmentation was used to compare cerebral blood flow as measured with ASL.

Results: In the CFS population, FA was increased in the right arcuate fasciculus ($P = .0015$), and in right-handers, FA was also increased in the right inferior longitudinal fasciculus (ILF) ($P = .0008$). In patients with CFS, right anterior arcuate FA increased with disease severity ($r = 0.649$, $P = .026$). Bilateral white matter volumes were reduced in CFS (mean $\pm$ standard deviation, 467.581 mm$^3 \pm 47.610$ for patients vs 504.864 mm$^3 \pm 68.126$ for control subjects, $P = .0026$), and cortical thickness increased in both right arcuate end points, the middle temporal ($T = 4.25$) and precentral ($T = 6.47$) gyri, and one right ILF end point, the occipital lobe ($T = 5.36$). ASL showed no significant differences.

Conclusion: Bilateral white matter atrophy is present in CFS. No differences in perfusion were noted. Right hemispheric increased FA may reflect degeneration of crossing fibers or strengthening of short-range fibers. Right anterior arcuate FA may serve as a biomarker for CFS.

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Online supplemental material is available for this article.
Chronic fatigue syndrome (CFS) is a debilitating disorder characterized by 6 or more months of persistent or relapsing fatigue without any associated medical or psychiatric disorder (1). The high prevalence of 2–4 per 1000 people in the United States (2,3), combined with the profound disability (4) and poor prognosis (5), motivates urgent scientific investigation. Brain imaging could aid in diagnosis and prognosis. However, structural imaging findings have been inconsistent: Three voxel-based morphometry studies demonstrated gray matter atrophy in differing locations when specified (6–8), while one study in which cerebrospinal fluid was segmented yielded no atrophy (9). Brain perfusion studies have shown inconsistent decreases (10), with a well-controlled study of monozygotic twins showing no differences (11). No technique has provided either a pathophysiological understanding of the disorder or served as a biomarker.

With diffusion-tensor imaging (DTI), the random motion of water is used to demonstrate brain microstructure and has been explored across a variety of neurodegenerative disorders. White matter microstructure may be altered in CFS as a consequence of inflammation (12) or as part of the neurocognitive physiology of fatigue (13), but it has not been investigated to date with DTI.

The purpose of this study was to (a) identify differences in gross brain structure in CFS by using T1-weighted gray and white matter volumetric analysis, (b) detect microstructural abnormalities underlying CFS by using DTI, and (c) detect global alterations in brain perfusion by using pseudocontinuous arterial spin labeling (ASL).

**Materials and Methods**

**Subjects**

This study began in June 2011, and follow-up was completed in November 2013. Institutional review board approval with a waiver of consent and a waiver of authorization were obtained, and a retrospective review was conducted from 2008 to the present for patients evaluated in the university CFS clinic. Inclusion criteria were (a) the clinical criteria for CFS (fatigue for a duration of 6 months or longer, along with having at least four of the eight Fukuda symptoms: impaired memory or concentration, sore throat, tender lymph nodes, headaches, muscle pain, joint pain, unrefreshing sleep, and postexertional malaise) and (b) ongoing memory and concentration symptoms that cause a severe enough impairment that the physician determined clinical magnetic resonance (MR) imaging was appropriate to exclude other diagnoses. All patients with CFS except one were evaluated by one clinician (J.G.M., with 10 years of experience with CFS). No other inclusion or exclusion criteria were used for patients with CFS. Among 259 charts reviewed, 15 patients were identified who met these criteria, and none of these patients were excluded. More than 300 control subjects from a database of volunteers were examined to find age-matched (within 1 year) and sex-matched participants without any history of major depression, CFS or chronic fatigue, or substance abuse in the past year; 28 eligible volunteers were contacted, and 14 chose to participate and underwent MR imaging. Control subjects provided informed consent in accordance with the institutional review board and the Health Insurance Portability and Accountability Act. The 20-item Multidimensional Fatigue Inventory (MFI-20) was administered to each subject, which has been validated as a reproducible instrument for the identification of CFS (14–17). The MFI-20 score is used to assess general, physical, and mental fatigue, as well as reduced motivation and activity; higher MFI-20 scores indicate increased severity. Subjects completed an expanded Edinburgh handedness inventory (http://www.brainmapping.org/shared/Edinburgh.php), with scores thresholded at 48.

**Implication for Patient Care**

- Right anterior arcuate FA may be a biomarker for CFS.
and higher. Four subjects declined this inventory and simply described themselves as left-handed, right-handed, or ambidextrous. Left-handed and ambidextrous subjects were pooled as non-right-handed subjects.

**MR Imaging Acquisition**

Subjects were imaged with one of two identical GE 3T 750HDx 60-cm–bore magnets (with scheduling dictating the imaging unit chosen) by using an eight-channel phased-array receive-only head coil and body–transmit coil. For volumetric assessment, we performed axial three-dimensional T1-weighted inversion-recovery spoiled gradient-echo and three-dimensional axial brain volume imaging with array spatial sensitivity encoding technique acceleration of 2, or ASSET 2, (repetition time msec/echo time msec, 9.15/3.7; inversion time msec, 450; flip angle, 13°; bandwidth, 25; field of view, 240 mm; 256 × 256 matrix; 1.2-mm-thick sections; 130 sections acquired; imaging time, 2 minutes 50 seconds). Two diffusion-weighted sequences were performed, the first with higher spatial resolution and the second with higher angular resolution: (a) one b of 0 mm/sec² image acquired, 40 directions at b of 2000 mm/sec², twice refocused, two-dimensional axial, ASSET 2, frequency right/left, 8000/95.5, field of view of 240 mm, 128 × 128 matrix reconstructed at 256 × 256, 2-mm-thick sections with 0-mm gap, 59 sections acquired, two signals acquired, and imaging time of 11 minutes; (b) and one b of 0 mm/sec² image acquired, 50 directions at b of 2000 mm/sec², twice-refocused two-dimensional axial, ASSET 2, frequency right/left, 8000/90.2, field of view of 240 mm, 96 × 96 matrix reconstructed at 256 × 256, 4-mm-thick sections with 0-mm gap, 36 sections acquired, two signals acquired, and imaging time of 13 minutes 44 seconds. Pseudocontinuous ASL was performed (three-dimensional axial imaging, 4800/10.6, bandwidth of 62, field of view of 220, 128 × 128 matrix, 4-mm-thick sections, 63 sections acquired, three signals acquired with one signal acquired for the unlabeled proton-density image, and imaging time of 4 minutes 50 seconds). One patient with CFS did not undergo ASL imaging because of a technical error.

**Segmentation: Volumetry**

A FreeSurfer pipeline analysis conducted by using the T1-weighted images (http://surfer.nmr.mgh.harvard.edu/) produced cortical, white matter, and subcortical segmentations (18). These segmentations underwent blinded manual editing (by J.K., with 2 years of neuroradiology experience, and M.M.Z., with 17 years of segmentation and quantitative neuroimaging experience) to improve skull stripping (typically, portions of the skull base, dura, and dural venous sinuses are erroneously characterized as cortex), extend the white matter segmentation to include all subcortical white matter (due to residual B0 inhomogeneity after correction), and retract the white matter segmentation when it seeps into the cortex (which occurs near primary somatosensory cortex at the vertex with its inherently reduced contrast with the subjacent white matter at T1-weighted imaging) (19). An additional step of blinded manual editing was performed for quantification of subcortical gray structures by using FreeView (by removing extra voxels from the hippocampal and thalamic segmentations). This produces a complete segmentation of the cortex, supratentorial white matter, deep gray nuclei, and ventricles. For computing total intracranial volume, we summed the supratentorial white matter, supratentorial white matter, and cerebrospinal fluid compartments (the latter was estimated from the sulcal, lateral, and third ventricular and choroid plexus volumes).

**DTI Acquisition**

The DTI acquisitions were imported separately, along with the T1-weighted volume, into two separate and independent instances of automated fiber quantification (http://white.stanford.edu/newlm/index.php/AFQ, performed by M.M.Z.) (20). Imaging volumes were inspected manually, and occasional artifactual diffusion-weighted imaging volumes were excluded from analysis. Data were corrected for motion with a rigid-body transformation and aligned to the b of 0 mm/sec² image, which was then aligned to the T1-weighted volume in anterior commissure–posterior commissure space. Automated fiber quantification performs automated tractography of 20 tracks, nine in each hemisphere (anterior thalamic radiations, corticospinal, cingulate cingulum, parahippocampal cingulum, inferior frontal occipital fasciculus, inferior longitudinal fasciculus [ILF], superior longitudinal fasciculus, uncinate fasciculus, and arcuate fasciculus) and two bilateral tracks (forceps major and minor of the corpus callosum). Each tract (a) was cleaned of extraneous fibers that deviated too far from the fiber core and truncated at standard region-of-interest (ROI) positions and (b) was normalized to 100 pixels in length. With this process, some tracks could be identified in all subjects.

**ASL Acquisition**

ASL signal intensity was divided by signal intensity in coplanar proton-density images and multiplied by a scaling factor to deliver maps of cerebral blood flow in milligrams per milliliter per minute (21) (www.nmr.mgh.harvard.edu/~jjchen/ASL.html). The proton-density image acquired as part of the ASL sequence was used to align with the T1-weighted volume by using the FMRIB (Functional MRI of the Brain) Software Library, or FSL, “flirt” function with six degrees of freedom and a normalized mutual information cost function, so cerebral blood flow images were in the same space as the FreeSurfer segmentations. This alignment was manually inspected by M.M.Z. to ensure accuracy. Cerebral blood flow was averaged over the segmentations of the cerebral cortex, supratentorial white matter, basal ganglia, thalamus, and hippocampi.

**Statistical Analysis**

All statistical analyses were performed by M.M.Z. For all volumetric comparisons, a regression was computed by using age, total intracranial volume,
disease, and handedness as independent variables, a recommended procedure for these types of analyses (22), by using Stata software (StataCorp, College Station, Tex). Uncorrected P values for the comparison of the volumes of the six regions evaluated—cortical gray matter, supratentorial white matter, thalami, hippocampi, basal ganglia, and rostral middle frontal gyri—are reported, along with the Bonferroni-corrected threshold for significance (.05/6 = .0083). The right and left white matter compartments and middle frontal gyri were evaluated separately by using the same regression procedure. Exclusively within the CFS population, a Pearson correlation coefficient was computed between regional volumes and MFI-20 scores.

For the cortical thickness evaluation, we first made global comparisons by examining the mean cortical thickness across both hemispheres together, again comparing via regression. We then used the FreeSurfer “qdec” module to compare cortical thickness between the CFS group and the control group, regressing age as a covariate (22), and handedness as an additional discrete factor. By using a different offset and different slope for each group and smoothing of 10 mm, we corrected for multiple comparisons with a false-discovery rate of .05. Cortical thickness was regressed in the CFS group with MFI-20 scores as an additional covariate, also by using “qdec.”

For the DTI, fractional anisotropy (FA) was sampled along each identified track and compared piecewise between groups by using an unpaired t test to compare both cohorts. Additional testing was performed in right-handed individuals from each group because differences in language lateralization can affect several tracks (including the right arcuate) (23). Automated fiber quantification uses a permutation-based method to correct for multiple comparisons between groups along a single track (24), producing a threshold P value equivalent to a corrected P value of .05. These P value thresholds vary per track, likely because tracks have unique correlation structures based on regional differences in subject anatomy, as well as differences attributable to crossing fibers. A track had a significant difference if the P value for the t test was below this threshold for at least one point along the track. The high-angular-resolution first DTI acquisition was used to generate a hypothesis: All tracks were tested for significant differences of FA, correcting for multiple comparisons within each track, but not across all tracks. The high-angular-resolution second DTI acquisition was used to test a hypothesis; Significant results from the hypothesis-generating data set were tested individually for significant differences in FA, correcting for multiple comparisons within each track, and not needing a correction across tracks because of the focused hypothesis testing. Maximal FA and mean axial diffusivity and radial diffusivity were computed over the anterior 10% of the right arcuate fasciculus. This maximal FA was correlated with the subject’s MFI-20 scores separately for patients and control subjects. Axial diffusivity and radial diffusivity were compared between groups by using an unpaired t test. A receiver operating characteristic (ROC) curve was plotted separately for the hypothesis-generating and hypothesis-testing data sets by identifying the accuracy of using maximal FA in the anterior 10% of the right arcuate fasciculus to diagnose CFS across multiple FA thresholds (with the FA threshold spaced .05 apart) by using Excel (Microsoft, Redmond, Wash). The 95% confidence intervals (CIs) for sensitivity and specificity were computed by using the Stata “diagnostic” module, and the area under the ROC curve was determined with the function “roctab.” For combined visualization of tractography and cortical thickness results, we first depicted the right arcuate and right ILF in one subject by using automated fiber quantification. Then, we drew ROIs around the three largest clusters of significantly different thicknesses by using “qdec” and transformed the ROIs into this single subject’s native space. To plot these ROIs alongside the tractography, the center of mass for each cluster was extracted by using FSL “fslstats” (25,26), and a spheroid was plotted with a volume equal to the cluster volume by using MATLAB software (Mathworks, Natick, Mass).

For the ASL, unpaired t tests were used to compare cerebral blood flow for each of the regions between groups, with P values reported uncorrected.

The results of all automated steps were verified by M.M.Z. either visually or numerically for plausibility. All results were interpreted with agreement by M.M.Z., A.L.R, M.M.R, J.K., and J.G.M.

Results

Clinical Data

Thirteen of the 15 patients with CFS and 10 of 14 control subjects were right-handed. Women constituted 55% of participants (eight patients with CFS and eight control subjects), and 45% of the participants were men (seven patients with CFS and six control subjects). Mean age ± standard deviation was 46.2 years ± 14.2 (range, 20–66 years) and was statistically equivalent for women and men (46.6 years ± 15.6 [range, 20–66 years] vs 45.6 years ± 12.9 [range, 23–60 years] for men; P = .86). Mean ages of the CFS and control groups were statistically equivalent (46.5 years ± 13.2 and 46.6 years ± 14.6, respectively; P = .98). MFI-20 scores were significantly different from patients compared with control subjects (MFI-20: 79.60 years ± 15.3 vs 27.64 years ± 8.54, respectively; P < .001). The mean duration of symptoms for CFS participants was 12.1 years ± 6.9.

Segmentation

Volumetry.—For the entire cohort, there was significantly lower total supratentorial white matter volume for patients with CFS compared with control subjects, when accounting for age, total intracranial volume, and handedness (Table 1). This reduction was present bilaterally (left, P = .0035; right, P = .0033). Additionally, total thalamic volume tended to be lower in CFS, but this did not reach significance after
correction for multiple comparisons. Total cortical gray matter volume was statistically equivalent. Prefrontal cortex volumes (identified in other studies of CFS, where foci are best matched by the rostral middle frontal gyr as segmented with FreeSurfer) (6,27) were also statistically equivalent between groups. However, the right middle frontal gyrus trended toward increased volume in the CFS group (15.446 mm³ ± 2771 in the CFS group and 14.721 mm³ ± 2158 in control subjects; P = .0212). Within the CFS population, the MFI-20 score trended toward a positive correlation with basal ganglia volume (r = 0.5564, P = .0312).

Thickness.—Global cortical thickness was equivalent between populations (2.437 mm ± 0.102 in patients with CFS vs 2.418 mm ± 0.091 in control subjects; P = .932) (Table 2, Fig 1). No focal left hemispheric differences were present. However, the right hemisphere demonstrated five regions of increased cortical thickness when accounting for age and handedness, which was most notable in the younger, not the older, patients with CFS, with no regions of decreased cortical thickness (Table 2, Fig 1). No regions demonstrated cortical thickness that increased significantly with age. No significant correlations with MFI-20 were present bilaterally in the CFS group.

**DTI in all subjects, right- and left-handed.**—In the high-resolution hypothesis-generating data set, the only significant difference was that the anterior right arcuate fasciculus (identified in 13 of 15 patients and 14 of 14 control subjects) demonstrated significantly higher FA in patients compared with control subjects, correcting for multiple-comparisons within but not across tracks (Table 3; Figs 2, 3). By performing the same unpaired t test just on the right arcuate fasciculus in the second hypothesis-testing DTI data set (also identified in 13 of 15 patients and 14 of 14 control subjects), we confirmed with a highly significant P value the same result by testing this single hypothesis (P = .0001, significant after correction within track). This FA increase was accompanied by an increase in axial diffusivity ([1.117 ± 0.089] × 10⁻³ mm²/sec for patients with CFS vs [1.005 ± 0.084] × 10⁻³ mm²/sec for control subjects, averaged over the anterior 10% of the arcuate fasciculus, pixel positions 90–100) and a decrease in radial diffusivity (0.364 ± 0.069 for patients vs 0.425 ± 0.039 for control subjects). The correlation between maximal FA over the same portions of this track and MFI-20 in the CFS population was not significant (r = 0.276, P = .4). Of note, the two end points of the right arcuate fasciculus are adjacent to the region of increased cortical thickening in the right precentral and middle temporal gyri (Figs 1, 3; Movie 1 [online]).

**DTI in right-handed subjects only.**—Right anterior arcuate FA was again significantly increased in CFS (Fig 2a, first DTI acquisition, P = .0006; Fig 2b, second DTI acquisition, P = .00036; both highly significant after correction within track). The correlation between maximal FA over the anterior 10% of this track and MFI-20 in the CFS population was significant in

### Table 1

<table>
<thead>
<tr>
<th>Brain Region</th>
<th>Volume in Patients with CFS (mm³)</th>
<th>Volume in Control Subjects (mm³)</th>
<th>β Value for the Fitted Slope</th>
<th>P Value for Disease Status in Regression Analysis</th>
<th>r Value for MFI-20 Scores in CFS</th>
<th>Bonferroni-corrected P Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cortical gray matter</td>
<td>448.899 ± 59.082</td>
<td>449.627 ± 47.163</td>
<td>13.120</td>
<td>0.0728</td>
<td>0.3658</td>
<td>0.1800</td>
</tr>
<tr>
<td>Supratentorial white matter</td>
<td>467.581 ± 47.610*</td>
<td>504.864 ± 68.126*</td>
<td>-17.439*</td>
<td>0.0026*</td>
<td>0.0214</td>
<td>0.0996</td>
</tr>
<tr>
<td>Thalami</td>
<td>13.115 ± 1870</td>
<td>14.430 ± 1812</td>
<td>-1.099</td>
<td>0.0469</td>
<td>0.2196</td>
<td>0.4317</td>
</tr>
<tr>
<td>Hippocampi</td>
<td>765.5 ± 528</td>
<td>7950 ± 781</td>
<td>-284</td>
<td>0.1341</td>
<td>0.1256</td>
<td>0.6555</td>
</tr>
<tr>
<td>Basal ganglia</td>
<td>19.982 ± 2951</td>
<td>19.321 ± 2832</td>
<td>72</td>
<td>0.8934</td>
<td>0.5564</td>
<td>0.0312</td>
</tr>
<tr>
<td>Rostral middle frontal gyri</td>
<td>30.323 ± 5471</td>
<td>30.144 ± 4733</td>
<td>1711</td>
<td>0.1357</td>
<td>0.2343</td>
<td>0.4006</td>
</tr>
</tbody>
</table>

Note.—r values demonstrate the correlations between CFS patient volumes and their corresponding MFI-20 scores. r values were calculated with the Pearson correlation coefficient and are shown with the associated P values, with a Bonferroni-corrected threshold of .0083 (to control for the six regions tested).

* Significant difference.

### Table 2

<table>
<thead>
<tr>
<th>Location</th>
<th>Area (mm²)</th>
<th>T Score</th>
<th>X Value</th>
<th>Y Value</th>
<th>Z Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lateral occipital</td>
<td>100.14</td>
<td>5.3592</td>
<td>-25.3</td>
<td>-88.4</td>
<td>-9</td>
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<tr>
<td>Precentral</td>
<td>89.19</td>
<td>5.3675</td>
<td>40.4</td>
<td>-10.7</td>
<td>34</td>
</tr>
<tr>
<td>Middle temporal</td>
<td>35.84</td>
<td>4.2466</td>
<td>83.3</td>
<td>-25.5</td>
<td>-12.3</td>
</tr>
<tr>
<td>Postcentral</td>
<td>16.84</td>
<td>4.5786</td>
<td>61.4</td>
<td>-11.9</td>
<td>17.2</td>
</tr>
<tr>
<td>Pars orbitalis</td>
<td>8.83</td>
<td>4.0795</td>
<td>40.2</td>
<td>48.4</td>
<td>-4.9</td>
</tr>
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</table>

Note.—X, Y, and Z refer to Montreal Neurological Institute 305 coordinates.
the hypothesis-generating data set but not the hypothesis-testing data set ($r = 0.649 \ [P = .026]$ and $r = 0.472 \ [P = .136]$, respectively). There was no correlation between MFI-20 and the same pixel locations in control subjects ($r = -0.02$). FA was also increased in the right anterior ILF (Figs 2c, 2d, 3; first DTI acquisition, $P = .0008$; second DTI acquisition, $P = .0018$; both significant after correcting within track). However, the peak FA was of greater magnitude difference and more consistent in location in the arcuate fasciculus compared with the ILF. The occipital lobe end point of the ILF was adjacent to the right occipital region of cortical thickening (Figs 1, 3; Movie 1 [online]). An ROC curve was calculated by using maximal FA in the anterior 10% of the right arcuate fasciculus to diagnose CFS in right-handed patients (Fig 4 shows the hypothesis-testing data set ROC). A threshold of 0.6 would be used to correctly diagnose CFS in right-handed patients in the hypothesis-generating data set, with a sensitivity, specificity, and area under the ROC curve of 72.7% (95% CI: 39.0%, 94.0%), 90.0% (95% CI: 55.5%, 99.7%), and 0.891 (95% CI: 0.696, 0.98) respectively.

**ASL Findings**

ASL demonstrated no significant difference in perfusion to the cortex (672 mL per 100 mg per minute ± 123 for patients with CFS vs 633 mL per 100 mg per minute ± 145 for control subjects; $P = .39$), supratentorial white matter (495 mL per 100 mg per minute ± 88 for patients with CFS vs 496 mL per 100 mg per minute ± 113 for control subjects; $P = .84$), basal ganglia (523 mL per 100 mg per minute ± 84 for patients with CFS vs 512 mL per 100 mg per minute ± 81 for control subjects; $P = .69$), thalami (599 mL per 100 mg per minute ± 116 for patients with CFS vs 587 mL per 100 mg per minute ± 116 for control subjects; $P = .78$), or hippocampi (587 mL per 100 mg per minute ± 106 for patients with CFS vs...
Radiology: Volume 274: Number 2—February 2015 • radiology.rsna.org 523

Table 3

<table>
<thead>
<tr>
<th>Track</th>
<th>All Patients</th>
<th>Right-Handers</th>
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<tbody>
<tr>
<td></td>
<td>P Value</td>
<td>P Value Threshold</td>
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<tr>
<td>Left thalamic radiation</td>
<td>.2144</td>
<td>.0027</td>
</tr>
<tr>
<td>Right thalamic radiation</td>
<td>.0360</td>
<td>.0019</td>
</tr>
<tr>
<td>Left corticospinal</td>
<td>.0087</td>
<td>.0033</td>
</tr>
<tr>
<td>Right corticospinal</td>
<td>.0499</td>
<td>.0024</td>
</tr>
<tr>
<td>Left cingulum cingulate</td>
<td>.0029</td>
<td>.0012</td>
</tr>
<tr>
<td>Right cingulum cingulate</td>
<td>.1325</td>
<td>.0019</td>
</tr>
<tr>
<td>Left cingulum hippocampus</td>
<td>.0566</td>
<td>.0021</td>
</tr>
<tr>
<td>Right cingulum hippocampus</td>
<td>.1107</td>
<td>.0027</td>
</tr>
<tr>
<td>Callosum forceps major</td>
<td>.0854</td>
<td>.0016</td>
</tr>
<tr>
<td>Callosum forceps minor</td>
<td>.1175</td>
<td>.0022</td>
</tr>
<tr>
<td>Left inferior fronto-occipital fasciculus</td>
<td>.0133</td>
<td>.0014</td>
</tr>
<tr>
<td>Right inferior fronto-occipital fasciculus</td>
<td>.0099</td>
<td>.0011</td>
</tr>
<tr>
<td>Left ILF</td>
<td>.0041</td>
<td>.0026</td>
</tr>
<tr>
<td>Right ILF</td>
<td>.0055</td>
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<tr>
<td>Left superior longitudinal fasciculus</td>
<td>.0842</td>
<td>.0040</td>
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<tr>
<td>Right superior longitudinal fasciculus</td>
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<td>.0045</td>
</tr>
<tr>
<td>Left uncinate</td>
<td>.2574</td>
<td>.0031</td>
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<tr>
<td>Right uncinate</td>
<td>.2390</td>
<td>.0051</td>
</tr>
<tr>
<td>Left arcuate</td>
<td>.0210</td>
<td>.0022</td>
</tr>
<tr>
<td>Right arcuate</td>
<td>.0015*</td>
<td>.0023</td>
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</table>

* P values are significant after correcting for multiple comparisons within the track (ie, under the P value threshold).

Discussion

This DTI study of CFS demonstrated increased FA in the right anterior arcuate fasciculus, and this increase correlated with disease severity. An automated, user-independent, microstructural DTI analysis identified this increased FA and verified it in a second data set from the same subjects analyzed independently. In right-handed patients with CFS, this increase was correlated with the patients’ MFI-20 scores. Thus, in populations of CFS that have severe concentration and memory problems, right anterior arcuate FA may serve as a biomarker for the disease. An analysis of volumetric data in the same subjects was used to identify reduced bilateral supratentorial white matter volumes, suggesting a global white matter process. Additionally, two cortical regions connected via the arcuate fasciculus exhibited increased thickness (28–30): the right middle temporal and precentral gyri. Similarly, in right-handers, the ILF showed that increased FA anteriorly increased thickness of a corresponding right occipital region. Our examination yielded no differences in perfusion.

While microstructure has not been studied to date in CFS, increased FA is an unexpected finding for a disorder characterized by reduced cognitive abilities. However, increased FA has been previously reported in the corona radiata in Alzheimer disease (31), possibly due to degeneration of crossing fibers. Given that the difference in CFS only involved the anterior 10% of the arcuate, we suspect this local phenomenon could represent strengthening of fibers that join a small portion of the anterior arcuate, or alternatively a weakening of crossing fibers. Consistent with our study, others have reported that the right arcuate fasciculus is sometimes not found with tractography, in part because the arcuate is more lateralized to the left hemisphere in right-handers (23,32). We found that arcuate differences between patients with CFS and control subjects were most striking among right-handed patients, suggesting that hemispheric differences with handedness and language (23) are an additional source of variance.

Our volumetry results stand in contradistinction to the literature and show reduced gray matter volumes, either globally or in the prefrontal cortices (6,7,27,33). In fact, we found that the prefrontal volumes trended toward being slightly higher in patients with CFS. Previous investigators have exclusively used voxel-based morphometry, which is based on many factors other than cortical...
Figure 2:  

(a) FA is plotted along the right arcuate fasciculus from the first diffusion-weighted acquisition; anterior = 0, posterior = 100. Thin blue dotted lines correspond to ±1 standard error. Only right-handers are included in this Figure. The blue arrow highlights the region of maximal increase of FA in patients with CFS compared with control subjects. 

(b) A plot is given for the second diffusion-weighted acquisition. 

(c, d) Similar plots are given for the right ILF, with the yellow arrow on c similarly highlighting the region of maximal increase of FA.

thickness (34). The discordance between the literature and our results may be explained by our precise methods, with a careful examination of cortical architecture and correction for covariates, such as age and total intracranial volume (22).

Although we did not control for caffeine exposure (35), our negative ASL result is consistent with a well-controlled perfusion study of CFS involving monozygotic twins (11), suggesting perfusion is not affected in CFS.

Even though the hypothesis-testing data set confirms the veracity of increased FA in the right anterior arcuate, it was determined in the same subjects and cannot be considered a fully independent data set. The correlation between MFI-20 and right anterior arcuate FA was only significant in the hypothesis-generating data set, not the hypothesis-testing data set. Although this may be technically related to the reduced spatial resolution of the latter data set, the utility of arcuate FA as a disease biomarker cannot be conclusively confirmed on this study. While the regions of increased cortical
thickness are in close proximity to the end points of the arcuate and ILF, these foci are relatively small and imperfectly aligned, suggesting a small effect size. Overall, this study has a small number of subjects, so all of the findings in this study require replication and exploration in a larger group of subjects.

Future work includes validating this finding in a larger right-handed cohort, teasing apart the components of crossing and short-range fibers in the anterior arcuate by using a multiple-shell (multiple b value) acquisition and examining the time-course in a longitudinal study, possibly with interventions, and investigating right-hemisphere networks with resting-state functional MR imaging.

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