White matter change with apathy and impulsivity in frontotemporal lobar degeneration syndromes

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Neurology® 2018;90:e1066-e1076. doi:10.1212/WNL.0000000000005175

Abstract

Objective
To identify the white matter correlates of apathy and impulsivity in the major syndromes associated with frontotemporal lobar degeneration, using diffusion-weighted imaging and data from the PiPPIN (Pick’s Disease and Progressive Supranuclear Palsy: Prevalence and Incidence) study. We included behavioral and language variants of frontotemporal dementia, corticobasal syndrome, and progressive supranuclear palsy.

Methods
Seventy patients and 30 controls underwent diffusion tensor imaging at 3-tesla after detailed assessment of apathy and impulsivity. We used tract-based spatial statistics of fractional anisotropy and mean diffusivity, correlating with 8 orthogonal dimensions of apathy and impulsivity derived from a principal component analysis of neuropsychological, behavioral, and questionnaire measures.

Results
Three components were associated with significant white matter tract abnormalities. Carer-rated change in everyday skills, self-care, and motivation correlated with widespread changes in dorsal frontoparietal and corticospinal tracts, while carer observations of impulsive–apathetic and challenging behaviors revealed disruption in ventral frontotemporal tracts. Objective neuropsychological tests of cognitive control, reflection impulsivity, and reward responsiveness were associated with focal changes in the right frontal lobe and presupplementary motor area. These changes were observed across clinical diagnostic groups, and were not restricted to the disorders for which diagnostic criteria include apathy and impulsivity.

Conclusion
The current study provides evidence of distinct structural network changes in white matter associated with different neurobehavioral components of apathy and impulsivity across the diverse spectrum of syndromes and pathologies associated with frontotemporal lobar degeneration.
Apathy and impulsivity are common and often coexistent in neurodegenerative disorders, including the clinical syndromes resulting from frontotemporal lobar degeneration (FTLD). They are difficult to treat and cause substantial patient morbidity and carer distress. Research into the causes and treatment of apathy and impulsivity is challenging because they are both multifaceted constructs: apathy reflects abnormal goal-directed behavior, from dysfunction in cognitive, emotional, and behavioral domains, while impulsivity is the tendency to act prematurely, without forethought or appropriate consideration of risk.

Resolving the neurobiological basis of apathy and impulsivity in neurodegenerative disease would facilitate the development and assessment of effective treatments and neuroprotective strategies. Herein, we focus on the heterogeneous clinical syndromes associated with FTLD, including behavioral variant frontotemporal dementia (bvFTD), primary progressive aphasias (nonfluent agrammatic variant [nvPPA], semantic variant [svPPA], and logopenic variant PPA), progressive supranuclear palsy (PSP), and the corticobasal syndrome (CBS).

We tested the hypothesis that across these diverse clinical syndromes, regionally specific pathology of white matter tracts as measured by diffusion tensor imaging (DTI) leads to different profiles of apathetic and impulsive behaviors. We consider the spectrum of FTLD disorders, rather than each separate syndrome, for 2 reasons. First, there is phenotypic overlap between syndromes. Second, apathy and impulsivity occur to a variable degree in each disorder, even where they are not diagnostic criteria. We predicted that separate dimensions of apathy and impulsivity would be associated with degeneration of distinct white matter tracts in neural systems supporting motivational and cognitive control.

Methods

Standard protocol approvals, registrations, and patient consents

The study was approved by the Cambridge 2 research ethics committee (reference 12/EE/0475) and supported by the National Institute for Health Research clinical research network (ID-15504). Informed consent was obtained at each study visit, with the personal consultee process used for participants who lacked mental capacity, in accordance with UK law.

Participants

The Pick’s Disease and Progressive Supranuclear Palsy: Prevalence and Incidence (PiPPIN) study recruited 204 participants. Recruitment and diagnostic criteria have been published previously. In brief, patients met clinical diagnostic criteria for behavioral and language variants of frontotemporal dementia (svPPA, nvPPA, logopenic variant PPA, and “other PPA” [not meeting criteria for 1 of the 3 defined subtypes]), CBS and possible, probable, or definite PSP (predominantly PSP Richardson syndrome under the revised criteria). Fifty healthy age- and sex-matched controls with no significant neurologic or psychiatric history were recruited. Participants were tested on their usual medication: 40% took “antidepressant” medications (for affective or behavioral indications), 29% dopaminergic medication, 4% antipsychotic medication, and 37% other centrally acting medications (benzodiazepines, antiepileptic, analgesics, pregabalin, or cholinesterase inhibitor). One hundred forty-nine patients and 50 controls underwent neuropsychological assessment, while advanced disease or death prevented assessment of the remaining patients.

One hundred participants underwent diffusion-weighted MRI. After quality control (excluding 1 patient and 2 controls), our imaging subset comprised 69 patients (22 PSP, 14 bvFTD, 14 CBS, 11 nvPPA, 4 svPPA, 4 other PPA) and 28 controls. To approximate group sizes, we evaluated PPA cases as a group. The scanned patients did not differ significantly from the nonscanned patients (table e-1, links.lww.com/WNL/A261).

Cognitive and behavioral assessments

The test battery examined the major components of apathy and impulsivity (table 1). Questionnaires sought multiple perspectives, including clinician, patient, and carer. Computerized behavioral tasks included measures of response inhibition (restraint: Go/NoGo, and cancellation: stop signal task), reflection impulsivity (information sampling task), and reward sensitivity (cued reinforcement reaction time task, Cambridge Gambling Task). Saccade and motor versions of the Go/NoGo task were used in view of the motor impairment inherent to some FTLD syndromes. We also assessed potential confounds including depression (Beck Depression Inventory–II), anhedonia (Snaith-Hamilton Pleasure Scale),...
<table>
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<tr>
<th>Measurement</th>
<th>Description</th>
<th>Variables for final PCA</th>
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</thead>
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<tr>
<td><strong>Questionnaires</strong></td>
<td></td>
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<tr>
<td>AES</td>
<td>18 items assessing emotional, behavioral, and cognitive constructs of apathy. All 3 available versions (patient, carer, clinician) were used.</td>
<td>AES 1: Patient ratings, AES 2: Carer and clinician</td>
</tr>
<tr>
<td>BIS</td>
<td>30-item self-report questionnaire reflecting the multifactorial structure of impulsivity. Outcome variables include attention, motor, self-control, cognitive complexity, perseverance, and cognitive instability subscores.</td>
<td>BIS 1: Attention, self-control, cognitive complexity, perseverance, BIS 2: Motor, cognitive instability</td>
</tr>
<tr>
<td>BIS/BAS</td>
<td>24-item self-report questionnaire based on Gray's biopsychological theory of personality. Outcome variables include the BIS subscore, reflecting aversive behaviors, and the BAS drive, fun-seeking and reward responsiveness subscores, reflecting appetitive behaviors.</td>
<td>BIS/BAS 1: BAS subscores, BIS/BAS 2: BIS subscore</td>
</tr>
<tr>
<td>MEI</td>
<td>27-item self-rated questionnaire developed to evaluate reductions in motivation and energy in depression research, although frequently used in other disease areas. The total score is the major outcome variable.</td>
<td>Total score</td>
</tr>
<tr>
<td>SHAPS</td>
<td>14-item self-rated questionnaire targeting hedonic capacity (anhedonia). The total score is the major outcome variable.</td>
<td>Total score</td>
</tr>
<tr>
<td>BDI-II</td>
<td>21-item self-rated depression questionnaire. The total score is the major outcome variable. The latest version, BDI-II, is designed for individuals aged 13 years and older. Cutoff scores are well established: 0–9: minimal depression; 10–18: mild depression; 19–29: moderate depression; and 30–63: severe depression.</td>
<td>Total score</td>
</tr>
<tr>
<td>CBI-R</td>
<td>45-item carer-rated questionnaire developed to evaluate behavioral changes associated with dementia. Outcome variables include memory/orientation, everyday skills, self-care, abnormal behavior, mood, beliefs, eating habits, sleep, stereotypical behavior, and motivation subscores.</td>
<td>CBI 1: Challenging behaviors, CBI 2: Everyday skills and self-care</td>
</tr>
<tr>
<td>NPI</td>
<td>12-item carer-rated questionnaire assessing the severity and distress caused by various behavioral disturbances. For the purposes of this study, only the apathy and disinhibition subscores were used.</td>
<td>NPI apathy/disinhibition subscores</td>
</tr>
<tr>
<td>Kirby</td>
<td>Serial forced-choice questionnaire to quantify the tendency to prefer small immediate rewards over larger delayed rewards. Outcome variables included the difference in K value, calculated as the difference in delayed discounting (K) from small to large delayed rewards (Klarge − Ksmall), and termed Kdiff.</td>
<td>Kdiff single score (note no difference to component structure if using standardized outcome measures [K]).</td>
</tr>
<tr>
<td><strong>Behavioral tasks</strong></td>
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<tr>
<td>CANTAB IST</td>
<td>Reflection impulsivity task administered on a touch-screen computer. Participants were presented with a 5 × 5 matrix of 25 gray boxes that, when selected, turned blue/yellow. On fixed trials (5), participants were instructed to open as many boxes as they liked, before deciding whether there were mostly blue or yellow boxes. On decreasing trials (5), every selected box subtracted 10 points from a starting 250, to encourage faster decision-making. Correct responses = 100 points; incorrect = −100 points. Outcome measures: probability of being correct, mean box-opening latency, mean color-decision latency, mean boxes opened per trial, sampling errors, discrimination errors, and total correct decisions.</td>
<td>IST 1: Proportion of correct trials, boxes opened, total correct, IST 2: Box and color latency, IST 3: Sampling error, boxes opened</td>
</tr>
<tr>
<td>CRRT</td>
<td>Reward sensitivity task measuring &quot;reinforcement-related speeding,&quot; administered on a laptop and 3-button press pad. Before each trial, participants observed a colored rectangle signaling the probability of reward following a correct response (20% vs 80% probability). Participants then identified the &quot;odd-one-out&quot; of 3 circles to receive feedback: 100 points for a fast correct, 1 point for a slow correct response, and 0 points for an incorrect response. Forty practice trials without feedback were used to titrate reaction time thresholds to individual differences in cognitive speed. Outcomes: speeding, total errors.</td>
<td>CRRT 1: Difference speeding, speeding FH, errors, CRRT 2: Difference speeding, speeding SH</td>
</tr>
</tbody>
</table>
and akinesia (PSP Rating Scale and reaction times), as discussed in Lansdall et al.\textsuperscript{3}\textsuperscript{,}3

SPSS version 22.0 (IBM Corp., Armonk, NY) was used for behavioral and neuropsychological analysis. Two-sample \textit{t} tests, corrected for multiple comparisons, were used for group comparisons. Principal component analysis (PCA) identified the major components of apathy and impulsivity.\textsuperscript{3} In brief, PCAs were run on control and patient data combined (n = 199; noting no major difference to the component structure using patient data only) with varimax rotation and mean replacement for missing data. The correlation matrix was used for component extraction based on Kaiser and Cattell criteria (whichever was more inclusive), while Kaiser-Meyer-Olkin and Bartlett test of sphericity confirmed the adequacy of the sample for PCA. Where questionnaires or tasks had multiple outcome measures, we ran a “local PCA.” A “final PCA” included the lead component loadings from local PCAs and total scores, accuracy (d-prime [d’]) or relevant subscores. Component scores were compared across groups using analysis of variance with post hoc least significant difference correction, and correlated with disease severity measures using Pearson correlations.

### Magnetic resonance imaging

Diffusion-weighted images were acquired using a Siemens Magnetom Tim Trio (Siemens, Erlangen, Germany) with a 63-direction gradient sequence with: b value 1,000 s/mm\textsuperscript{2}; repetition time 7,800 milliseconds; echo time 90 milliseconds; axial in-plane acquisition matrix 96 × 96; field of view 192 × 192 mm; slice thickness 2 mm; and a total of 63 contiguous slices with in-plane resolution 2-mm isotropic. An additional b value of 0 s/mm\textsuperscript{2} image was acquired.

Images were processed using FMRIB Software Library (FSL version 5.0; www.fmrib.ox.ac.uk/fsl), correcting for eddy currents and participant motion by affine registration to the first b0 image (FSL eddy_correct); bvecs were rotated (fdt_rotate_bvecs). The b0 image was extracted and a brain mask created (Brain Extraction Tool). Diffusion tensors were fitted (dift) to create maps of fractional anisotropy (FA) and mean diffusivity (MD). FA maps from 5 participants from each group were nonlinearly registered to the

### Table 1

<table>
<thead>
<tr>
<th>Measurement</th>
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<th>Variables for final PCA</th>
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</thead>
<tbody>
<tr>
<td>CANTAB SST</td>
<td>A response inhibition task (action cancellation), administered on a touch screen and 2-button press pad. Stimuli were presented on a computer screen and participants were instructed to press the right/left button as quickly as possible in response to the right/left arrow. For the test trials (64), participants were instructed to refrain from responding when they heard an auditory signal (beep), presented in 25% of trials (randomly dispersed). The delay between presentation of the arrow stimuli and the stop signal (stop signal delay) varied, in order to estimate the stop signal reaction time (time taken to successfully inhibit a response). The major outcome variables included SSD, SSRT, total correct responses, direction errors, and mean/median reaction times for all Go trials.</td>
<td>SST: SSRT, correct responses (proportion of successful stops), median reaction time on Go trials</td>
</tr>
<tr>
<td>Saccade NoGo</td>
<td>The saccadic NoGo task used direct infrared oculography from a head-mounted saccadometer (Ober Consulting Poland). Each session included 300 trials, following 10 calibration trials. Participants fixated centrally (red/green dots) on a screen at approximately 1.5-m distance. After 300 milliseconds, one of the central cues was removed and a red dot was presented at −10° or +10° horizontal displacement (randomized, 50:50). In 50% of trials, the green central cue remained and participants responded by a saccade to lateral target (Go trials). In NoGo trials, the red central cue remained and participants refrained from making a saccade. Outcome variables: calculated d’.</td>
<td>d’</td>
</tr>
<tr>
<td>Motor NoGo</td>
<td>The motor NoGo task was analogous to the saccadic task but used a joystick operated by the right hand (see supplementary material for details). Outcome measures for NoGo tasks included d’ for performance accuracy, commission and omission error rates, and reaction times. Calculated d’: Lower values reflect decreased “hits” (correct on Go trials) and increased false alarms (Go on NoGo trials: commission errors).</td>
<td>d’</td>
</tr>
<tr>
<td>CGT CANTAB</td>
<td>Participants were presented with a row of red and blue boxes and were instructed to guess which color box a yellow token was placed under, responding by touching the boxes containing the words “red” or “blue.” In the gambling stages, participants started with 250 points and could select their decision confidence by gambling a certain proportion of these points, which were displayed in either ascending (part 1) or descending (part 2) order. Participants were instructed to obtain as many points as possible, and the total accumulated points were displayed on the screen throughout. The gambling task was removed after 37 patients because of difficult task engagement, even following simplification of the task.</td>
<td>NA</td>
</tr>
</tbody>
</table>
FMRIB58_FA_1 mm target (tbss_2_reg). The warped FA images were averaged to produce a study-specific FA template.\textsuperscript{17} Registration was repeated for all participants using this study-specific FA template as target, bringing all participants into the same anatomical space. From the study-specific template, a mean FA skeleton was produced, and individual FA skeletons were mapped to it (threshold = 0.2). The transformations putting the individual FA maps into the skeletonized standard space were applied to MD maps.

Tract-based spatial statistics were used to examine the relationships between changes in diffusion metrics and behavior.\textsuperscript{18} Correlations between the skeleton DTI tracts and components of apathy and impulsivity were assessed by nonparametric permutation analysis using FSL randomise with threshold-free cluster enhancement (TFCE) correction, 2-dimensional optimization, and 5,000 permutations. The design matrix contained a constant term to model the intercept and each of the 8 orthogonal principal components of behavior. Cluster significance was tested at $p < 0.01$ and $p < 0.05$, corrected for multiple comparisons. White matter was labeled using the JHU (Johns Hopkins University) white-matter tractography atlas and ICBM-DTI-81 (International Consortium of Brain Mapping) white-matter labels atlas.

Results

Neuropsychological and behavioral results

Demographic, cognitive, neuropsychological, and behavioral results of patients and control participants who underwent DTI are displayed in table 2. Groups were matched for age and sex, while patients were impaired in cognition, disease severity, and most measures of apathy and impulsivity.

The PCA identified 8 components (table 3; table e-2, links.lww.com/WNL/A261).\textsuperscript{3} Short summary terms were assigned to each according to their major loadings, after Lansdall et al.\textsuperscript{3} Component 1, termed “patient-rated change,” reflected self-ratings of apathy (Apathy Evaluation Scale [AES]), impulsivity (Barratt Impulsiveness Scale), anhedonia (Snaith-Hamilton Pleasure Scale), depression (Beck Depression Inventory–II), and motivation (Motivation and Energy Inventory). Components 2 and 3 were carer-based, weighted toward the AES, Cambridge Behavioral Inventory (CBI), and Neuropsychiatric Inventory (NPI); component 2, “carer-rated change in everyday skills/self-care,” reflected apathy (NPI apathy and AES), everyday skills, self-care, sleep, and motivation (CBI), while component 3, “carer-rated change in complex behaviors,” reflected apathy (AES), impulsivity (NPI disinhibition), and stereotypic/complex behaviors (CBI). Performance on the Go/NoGo, information sampling, and cued reinforcement tasks loaded onto component 4, termed “impulsive behavior.” Kaiser-Meyer-Olkin statistic = 0.743 and Bartlett test $\chi^2_{128} = 508; p < 0.001$ confirmed data suitability for PCA. Patient-rated questionnaires, carer-rated questionnaires, and objective behavioral measures loaded onto distinct components, including positive weighting of both apathy and impulsivity measures.

Apathy and impulsivity were observed across the spectrum of clinical syndromes, reflecting their transdiagnostic nature. Significant differences between diagnostic groups were observed for loadings on components 1–4 (figure 1).

Diffusion tensor imaging

Tract-based spatial statistics identified significant (TFCE-corrected $p < 0.01$) changes in white matter in relation to carer-rated change in everyday skills and self-care (component 2, yellow-red) and carer-rated change in complex behaviors (component 3, blue-green; $p < 0.01$) (figure 2). Changes in MD and FA were complementary and highlighted concordant patterns of white matter change in relation to carer-rated change in everyday skills and self-care (component 2) and carer-rated change in complex behaviors (component 3). Loss of everyday skills correlated with FA (negative) and MD (positive) in the genu, body, and splenium of the corpus callosum, anterior and posterior corona radiata, corticospinal tracts, and posterior thalamic radiation (table e-3, links.lww.com/WNL/A261; figures e-1–e-3, links.lww.com/WNL/A260). Complex behaviors, including impulsivity, correlated with FA (negative) and MD (positive) in frontal-temporal connections between the orbital- and ventrolateral-prefrontal cortex, anterior cingulate, and temporal pole, including the genu and body of the corpus callosum, anterior limb of the internal capsule, anterior thalamic radiation, and anterior corona radiata (figures e-4 and e-5). The anterior–posterior dissociation between components 3 and 2 is most apparent for MD (figure 2A). The longitudinal, fronto-occipital and uncinate fasciculi, and forceps major and minor were associated with both carer-rated components, with a more restricted (anterior) distribution in relation to complex behaviors (figures 2, e-4, and e-5). At the more liberal threshold of $p < 0.05$ (TFCE-corrected), component 4 correlated with MD changes in regions connecting the presupplementary motor area and dorsolateral prefrontal cortex, and occipital lobe (thalamic radiation, forceps major, inferior fronto-occipital fasciculus, and inferior longitudinal fasciculus; figures e-1 and e-6).

Discussion

Distinct spatial distributions of white matter pathology are related to separate dimensions of apathy and impulsivity, across multiple syndromes associated with FTLD. Carers’ ratings of complex and challenging behaviors (including apathy and impulsivity, component 3) were associated with anterior changes in the white matter tracts connecting ventrolateral and orbitofrontal cortex and temporal poles. In contrast, carers’ ratings of everyday skills, self-care, and apathy correlated with changes in frontal, parietal, and corticospinal tracts. Our data also show that (1) apathy and impulsivity are positively correlated, and (2) they are present in all syndromes associated with FTLD. These critical results reinforce the phenotypic overlap between disorders, reflected in new diagnostic terms such as PSP-CBS, PSP-F...
Table 2  Demographics and neuropsychiatric and behavioral results for imaged patients and controls

<table>
<thead>
<tr>
<th>Variable</th>
<th>Imaged controls</th>
<th>Imaged patients</th>
<th>t Test, p value</th>
<th>PSP</th>
<th>CBS</th>
<th>PPA</th>
<th>bvFTD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographics and cognition/ function</td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>No.</td>
<td>28</td>
<td>69</td>
<td>NA</td>
<td>22</td>
<td>14</td>
<td>19</td>
<td>14</td>
</tr>
<tr>
<td>Age, y</td>
<td>68.4 ± 6.0</td>
<td>68.7 ± 8.0</td>
<td>NS</td>
<td>71.4 ± 7.4</td>
<td>66.9 ± 8.0</td>
<td>71.2 ± 7.5</td>
<td>63.9 ± 7.4</td>
</tr>
<tr>
<td>Sex, M/F</td>
<td>15/13</td>
<td>38/31</td>
<td>NS</td>
<td>12/10</td>
<td>7/7</td>
<td>11/8</td>
<td>8/6</td>
</tr>
<tr>
<td>ACE-R total (/100)</td>
<td></td>
<td></td>
<td></td>
<td>78.6 ± 11.8</td>
<td>66.1 ± 25.3</td>
<td>53.4 ± 21.9</td>
<td>67.2 ± 25.1</td>
</tr>
<tr>
<td>MMSE total (/30)</td>
<td></td>
<td></td>
<td></td>
<td>25.9 ± 4.3</td>
<td>21.7 ± 8.2</td>
<td>19.8 ± 7.3</td>
<td>23.5 ± 6.7</td>
</tr>
<tr>
<td>FRS % score (/100)</td>
<td></td>
<td></td>
<td></td>
<td>44.4 ± 29.2</td>
<td>34.5 ± 25.7</td>
<td>51.5 ± 29.7</td>
<td>26.0 ± 12.3</td>
</tr>
<tr>
<td>PSP-RS</td>
<td>NA</td>
<td>29.9 ± 18.6</td>
<td>NS</td>
<td>40.0 ± 11.4</td>
<td>37.3 ± 19.0</td>
<td>7.3 ± 5.4</td>
<td>16.0 ± 10.5</td>
</tr>
<tr>
<td>FAB</td>
<td>17.2 ± 0.9</td>
<td>10.5 ± 4.2</td>
<td>NS</td>
<td>11.4 ± 3.4</td>
<td>10.8 ± 4.8</td>
<td>8.9 ± 3.8</td>
<td>10.9 ± 5.1</td>
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<tr>
<td>Questionnaires</td>
<td></td>
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<tr>
<td>AES (/72)</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Careerb</td>
<td>24.3 ± 5.4</td>
<td>46.9 ± 12.4</td>
<td>NS</td>
<td>47.2 ± 11.1</td>
<td>47.4 ± 10.7</td>
<td>41.2 ± 14.9</td>
<td>53.6 ± 9.4</td>
</tr>
<tr>
<td>Patientc</td>
<td>24.5 ± 5.2</td>
<td>36.7 ± 9.2</td>
<td>NS</td>
<td>39.7 ± 10.9</td>
<td>35.2 ± 5.7</td>
<td>37.6 ± 6.3</td>
<td>32.6 ± 10.2</td>
</tr>
<tr>
<td>Clinicianc</td>
<td>25.4 ± 7.6</td>
<td>43.4 ± 9.6</td>
<td>NS</td>
<td>46.6 ± 10.8</td>
<td>42.4 ± 8.4</td>
<td>38.8 ± 10.0</td>
<td>43.4 ± 6.7</td>
</tr>
<tr>
<td>BIS (/120)</td>
<td>57.1 ± 7.8</td>
<td>64.2 ± 7.8</td>
<td>NS</td>
<td>65.4 ± 7.7</td>
<td>61.1 ± 10.5</td>
<td>65.4 ± 7.0</td>
<td>63.7 ± 6.3</td>
</tr>
<tr>
<td>BIS/BASc</td>
<td></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>BIS subscore</td>
<td>20.3 ± 3.0</td>
<td>20.8 ± 4.7</td>
<td>NS</td>
<td>19.9 ± 3.3</td>
<td>21.9 ± 3.0</td>
<td>22.4 ± 7.6</td>
<td>19.7 ± 3.3</td>
</tr>
<tr>
<td>BAS drive</td>
<td>10.5 ± 1.6</td>
<td>11.0 ± 3.3</td>
<td>NS</td>
<td>11.1 ± 3.1</td>
<td>9.5 ± 3.2</td>
<td>10.6 ± 3.2</td>
<td>12.7 ± 3.4</td>
</tr>
<tr>
<td>BAS fun-seeking</td>
<td>10.9 ± 2.1</td>
<td>11.2 ± 2.9</td>
<td>NS</td>
<td>10.7 ± 2.8</td>
<td>9.5 ± 3.1</td>
<td>11.8 ± 2.3</td>
<td>13.0 ± 2.6</td>
</tr>
<tr>
<td>BAS reward responsiveness</td>
<td>15.7 ± 2.8</td>
<td>16.4 ± 2.7</td>
<td>NS</td>
<td>16.1 ± 2.9</td>
<td>16.6 ± 2.2</td>
<td>16.4 ± 2.3</td>
<td>16.9 ± 3.2</td>
</tr>
<tr>
<td>MEI (/144)d</td>
<td>112.8 ± 15.8</td>
<td>80.3 ± 27.4</td>
<td>NS</td>
<td>67.5 ± 30.4</td>
<td>76.9 ± 25.6</td>
<td>86.7 ± 14.6</td>
<td>97.3 ± 24.9</td>
</tr>
<tr>
<td>BDI (/63)c</td>
<td>3.6 ± 4.1</td>
<td>13.3 ± 10.7</td>
<td>NS</td>
<td>19.0 ± 12.5</td>
<td>12.6 ± 8.2</td>
<td>9.0 ± 10.0</td>
<td>9.2 ± 6.0</td>
</tr>
<tr>
<td>SHAPS (/56)d</td>
<td>18.7 ± 4.8</td>
<td>22.4 ± 5.1</td>
<td>NS</td>
<td>22.4 ± 4.7</td>
<td>23.1 ± 5.7</td>
<td>20.6 ± 3.8</td>
<td>23.5 ± 6.3</td>
</tr>
<tr>
<td>NPI, fraction with positive responseb</td>
<td></td>
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</tr>
<tr>
<td>Apathy subscore</td>
<td>0.00 ± 0.00</td>
<td>0.60 ± 0.49</td>
<td>NS</td>
<td>0.60 ± 0.50</td>
<td>0.71 ± 0.47</td>
<td>0.42 ± 0.51</td>
<td>0.71 ± 0.47</td>
</tr>
<tr>
<td>Disinhibition subscore</td>
<td>0.04 ± 0.19</td>
<td>0.36 ± 0.48</td>
<td>NS</td>
<td>0.29 ± 0.46</td>
<td>0.14 ± 0.36</td>
<td>0.32 ± 0.51</td>
<td>0.77 ± 0.44</td>
</tr>
<tr>
<td>CBI-R (/180)b</td>
<td>4.5 ± 4.2</td>
<td>62.8 ± 35.2</td>
<td>NS</td>
<td>50.9 ± 33.9</td>
<td>69.8 ± 36.1</td>
<td>53.3 ± 37.8</td>
<td>85.2 ± 20.4</td>
</tr>
<tr>
<td>Kirby (difference)d</td>
<td>0.01 ± 0.02</td>
<td>0.01 ± 0.05</td>
<td>NS</td>
<td>0.03 ± 0.04</td>
<td>0.02 ± 0.05</td>
<td>0.01 ± 0.03</td>
<td>-0.001 ± 0.08</td>
</tr>
<tr>
<td>Behavioral tasks</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ISTd</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Probability of being correct, fixed</td>
<td>0.78 ± 0.10</td>
<td>0.75 ± 0.15</td>
<td>NS</td>
<td>0.68 ± 0.15</td>
<td>0.59 ± 0.24</td>
<td>0.64 ± 0.11</td>
<td>0.73 ± 0.19</td>
</tr>
<tr>
<td>Probability of being correct, decreasing</td>
<td>0.85 ± 0.12</td>
<td>0.67 ± 0.17</td>
<td>NS</td>
<td>0.75 ± 0.15</td>
<td>0.72 ± 0.14</td>
<td>0.68 ± 0.12</td>
<td>0.83 ± 0.14</td>
</tr>
<tr>
<td>CRRTd</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total errors</td>
<td>3.1 ± 2.9</td>
<td>4.2 ± 5.0</td>
<td>NS</td>
<td>3.7 ± 3.3</td>
<td>5.2 ± 5.0</td>
<td>7.0 ± 9.4</td>
<td>2.6 ± 2.1</td>
</tr>
</tbody>
</table>

Continued
language variants, especially svPPA, cause significant behavioral change including apathy and impulsivity. It also highlights the advantages of a dimensional approach that accommodates commonalities across groups and the convergence of syndromes with disease progression. In doing so, we confirm that even the language variants, especially svPPA, cause significant behavioral change including apathy and impulsivity.\(^1,3\)

The white matter abnormalities associated with challenging behaviors (component 3: AES, NPI disinhibition, and CBI abnormal/stereotypic behaviors, eating habits, and motivation) are consistent with previous studies linking apathy and impulsivity to abnormal white matter and metabolism in frontotemporal regions.\(^1,2,4\) They mirror white matter tract abnormalities in bvFTD,\(^7,19\) for which apathy and impulsivity are diagnostic criteria. Moreover, the uncinate fasciculus is linked to inhibitory control in bvFTD and apathy in Alzheimer disease,\(^17\) small vessel disease,\(^20\) PSP,\(^21\) and bvFTD.\(^1\) This suggests a common neural pathway, across disorders.

Carer-rated change in everyday skills, self-care, motivation, and apathy correlated with widespread white matter changes in the corpus callosum, corona radiata, superior longitudinal fasciculus, and thalamic radiation. In contrast to carer-rated change in complex behaviors, there was less emphasis on rostral frontotemporal change. PSP and CBS groups scored most highly on this component, although all groups scored higher on average than controls (figure 1). The results support previous volumetric analyses showing the following: (1) PSP degeneration of the brainstem and association and commissural fibers including superior cerebellar peduncles, corpus callosum, inferior longitudinal fasciculus, and superior longitudinal fasciculus;\(^21\) (2) CBS changes in frontoparietal tracts and corpus callosum;\(^22\) and (3) FTD widespread changes.\(^23\)

The widespread abnormalities are consistent with network-based disruption in FTLD,\(^24–26\) affecting broadly distributed frontotemporal networks rather than focal areas of damage. However, multifocal changes associated with carer reports may also reflect an inability to differentiate behavioral profiles using these questionnaires. Nonetheless, the tract-based statistics were broadly consistent with volumetric\(^1\) evidence of the breakdown of frontostriatal and frontotemporal circuits for motivation,\(^3,5\) coordinating the multiple cognitive domains necessary for planning and executing effective goal-directed behavior.\(^27\)

One difference between the former volumetric study and current DTI results is the absence of a tract-based deficit in relation to patients’ observations of their own symptomatology (table 3).\(^2\) There are several explanations for this discordance. First, patient ratings may reflect heterogeneous, multifocal changes in white matter, which prevent the identification of consistently localized tract correlates. Second, volumetric and DTI analyses assess fundamentally distinct neuropathologic features (tissue loss and T1 signal change vs the diffusion integrity of white matter connections), leading to different statistical associations. For example, patient ratings may reflect volumetric changes in deep white matter structures that are not captured by DTI. Third, the difference may reflect the limitations of white matter voxel-based morphometry,\(^18\) arising from normalization errors, mislocalization, or the partial-volume effects of smoothing, which can give rise to false-positives. The current tract-based method is less vulnerable to these issues, although there are limitations to the interpretation of DTI, which are discussed below. With the tract-based method, current white matter changes appear more extensive than the previously reported gray matter atrophy. For example, performance on the objective behavioral tasks correlated with white matter tract measures in the right frontal cortex, as well as white matter tracts near the regions of posterior and subcortical atrophy.\(^3\) This difference may be attributable to differential signal-to-noise of the 2 methods but may also reflect the core white matter pathophysiology in syndromes associated with FTLD.\(^7\).

### Table 2 Demographics and neuropsychiatric and behavioral results for imaged patients and controls (continued)

<table>
<thead>
<tr>
<th>Variable</th>
<th>Imaged controls</th>
<th>Imaged patients</th>
<th>t Test, p value</th>
<th>PSP</th>
<th>CBS</th>
<th>PPA</th>
<th>bvFTD</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSTd</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>SSRT</td>
<td>175.8 ± 42.8</td>
<td>447.0 ± 244.3</td>
<td>— ^a</td>
<td>449.4 ± 189.0</td>
<td>544.3 ± 430.7</td>
<td>471.8 ± 242.5</td>
<td>353.0 ± 152.2</td>
</tr>
<tr>
<td>Motor Go/NoGo d ^d</td>
<td>4.5 ± 0.3</td>
<td>3.2 ± 1.3</td>
<td>— ^a</td>
<td>3.4 ± 1.0</td>
<td>2.9 ± 1.6</td>
<td>3.0 ± 1.4</td>
<td>3.6 ± 1.5</td>
</tr>
<tr>
<td>Saccade d ^d</td>
<td>2.6 ± 0.9</td>
<td>0.8 ± 1.1</td>
<td>— ^a</td>
<td>0.7 ± 0.9</td>
<td>1.0 ± 0.8</td>
<td>0.5 ± 1.2</td>
<td>1.1 ± 1.4</td>
</tr>
</tbody>
</table>

Abbreviations: ACE-R = Addenbrooke’s Cognitive Examination—Revised; AES = Apathy Evaluation Scale; BAS = Behavioral Activation System; BDI = Beck Depression Inventory; BIS = Barratt Impulsiveness Scale; bvFTD = behavioral variant frontotemporal dementia; CBS = Cambridge Behavioral Inventory—Revised; CBS = corticobasal syndrome; CRRT = cued reinforcement reaction time; d = d-prime; FAB = frontal assessment battery; FRS = Frontotemporal Dementia Rating Scale; IST = Information sampling task; MEI = Motivation and Energy Inventory; MMSE = Mini-Mental State Examination; NA = not applicable; NPI = Neuropsychiatric Inventory; NS = not significant; PPA = primary progressive aphasia; PSP = progressive supranuclear palsy; PSP-RS = PSP Rating Scale; SHAPS = Snith-Hamilton Pleasure Scale; SST = stop signal task. Stats indicate Student t test results comparing imaged controls (n = 28) and patients (n = 67). ^a p < 0.01 (survives Bonferroni correction for multiple comparisons).  ^b Up to 5 missing data points.  ^c Up to 10 missing data points.  ^d More than 10 missing data points.  ^e p < 0.05.
Although the behavioral task performance showed weaker correlations with MD and the carer ratings, its anatomical correlates are of particular relevance. First, all patient groups performed worse than controls (figure 1D), confirming the objective neuropsychological deficits as a transdiagnostic phenomenon. Second, these regions (presupplementary motor area, dorsolateral prefrontal cortex, and inferior frontal gyrus; figure e-1, links.lww.com/WNL/A260) and their interconnections are strongly associated with cognitive and motor control in preclinical models and human studies. Reduced connectivity among these regions affects response inhibition and choices between alternate actions.32,33

Carer ratings and behavioral task performance all correlated with cognitive and functional decline. Previous studies have reported a link between apathy and poor outcome, with rapid cognitive and functional deterioration in apathetic patients compared to nonapathetic and depressed individuals.34 Further investigations assessing the prognostic implications of apathy and impulsivity in FTLD syndromes are warranted.

There are limitations to this study and caveats to the methods. DTI is an indirect measure of the physical properties of brain parenchyma, including white matter axon density, caliber, and myelination.35 The pathologic causes of abnormal diffusion are not fully elucidated. Even though the semiquantitative in vivo measures provide important anatomical insights, cross-validation with neuropathology is sparse. For example, preclinical studies link FA to myelination, membrane permeability, and fiber density in white matter.36 Comparative studies of anatomy across species and in FTLD post mortem are required to determine the pathologic mechanisms of the imaging changes we observe. Although different DTI metrics may reflect distinct processes (demyelination, neurodegeneration, gliosis, calcification, axonal degeneration, etc.), linking them to specific leucopathologies remains challenging. One must also consider artifacts from motion and registration errors, as multiple directional measurements are obtained at each voxel, introducing false-positive differences if movement differs by group.37 Registration poses significant challenges for FTLD groups with highly atrophic brains, obscuring some tracts and affecting the absolute
Figure 1 Component scores by diagnostic group

(A–D) Boxplots of principal component scores (2–4) by diagnosis for the imaged subset (n = 97). Bars indicate significant differences between each group and controls using analysis of variance with post hoc least significant difference tests (solid lines p < 0.001, dashed lines p < 0.05), and circles/stars represent outliers (1.5*IQR/3*IQR, respectively). (A) Component 1 representing patient-rated behavioral change as measured by the AES, Barratt Impulsiveness Scale, Snaith-Hamilton Pleasure Scale, Beck Depression Inventory–II, and Motivation and Energy Inventory. (B) Component 2 reflecting carer-rated change in everyday skills, self-care, and motivation as measured by the CBI subscores, AES, and NPI apathy subscore. (C) Component 3 reflecting carer-rated change in complex behaviors as measured by the CBI abnormal/stereotypic behaviors, eating habits, mood and motivation subscores, AES, and NPI disinhibition subscore. (D) Component 4 indicating poor performance on behavioral tasks of response inhibition (Go/NoGo motor and saccade), reflection impulsivity (information sampling task), and reward responsiveness (cured reinforcement reaction time task). Significant differences were also observed between groups for component 1 (F(4,92) = 7.462, p < 0.001, post hoc control vs PSP p < 0.001, vs PPA p < 0.05, PSP vs PPA p < 0.05, vs bvFTD p < 0.05), component 2 (F(4,92) = 9.132, p < 0.001, post hoc control vs PSP p < 0.001, vs CBS p < 0.001, PSP vs CBS p < 0.001, vs PPA, p < 0.05, CBS vs PPA p < 0.001, CBS vs bvFTD p < 0.001, PPA vs bvFTD p < 0.001), component 3 (F(4,92) = 23.832, p < 0.001, post hoc control vs bvFTD p < 0.001, vs PPA p < 0.05, CBS vs bvFTD p < 0.001, CBS vs PPA p < 0.001, PPA vs bvFTD p < 0.001), component 4 (F(4,92) = 10.902, p < 0.001, post hoc control vs PSP p = 0.001, CBS p < 0.05, PPA p < 0.001, bvFTD p < 0.001, PPA vs PSP p < 0.001, CBS vs PPA p < 0.001, CBS vs bvFTD p < 0.001, PPA vs bvFTD p < 0.001).

(E) Components 1–4 correlated with measures of cognition (ACE-R, MMSE, FAB) and disease severity (FRS, PSP-RS) with higher component scores reflecting greater cognitive impairment, functional decline, and disease severity (note Pearson correlation, p < 0.001 uncorrected here approximates p < 0.05 corrected for multiple comparisons). * = p < 0.05; ** = p < 0.001 unc; ACE-R = Addenbrooke’s Cognitive Examination–Revised; AES = Apathy Evaluation Scale; bvFTD = behavioral variant frontotemporal dementia; CBI = Cambridge Behavioral Inventory; CBS = corticobasal syndrome; FAB = frontal assessment battery; FRS = Frontotemporal Dementia Rating Scale; IQR = interquartile range; MMSE = Mini-Mental State Examination; NPI = Neuropsychiatric Inventory; PPA = primary progressive aphasia (all groups); PSP = progressive supranuclear palsy; PSP-RS = PSP Rating Scale.
diffusivities or eigenvalues. White matter change in areas with substantial gray matter atrophy may lead to changes in estimated FA/MD that reflect differences in the relative amounts of tissue types rather than change in white matter. In addition to these general DTI considerations, tract-based spatial statistics also has caveats. It attempts to overcome misregistration issues and ensure the same region (or voxel) corresponds across groups by creating a mean FA skeleton, onto which each individual’s FA is projected prior to statistics. This relies on accurate coregistration of FA images. White matter lesions that reduce FA may also alter the values chosen to represent the core of the tract during FA projection. Despite these risks, we favor tract-based spatial statistics over other frequently used whole-brain methods because of its increased sensitivity and power.

A neuropsychological battery is necessarily selective, and our findings are limited to the patients studied and the dimensions of apathy and impulsivity accessible to our tests and questionnaires. PiPPIN aimed to assess the multifaceted constructs of apathy and impulsivity, while accommodating the frailty of patients. Nonetheless, many patients found the Cambridge Gambling Task difficult to perform adequately. However, pathologic gambling is uncommon in FTLD disorders, and including this task in a subsidiary PCA did not alter the factor structure. Alternative tasks and questionnaires (e.g., cued reinforcement reaction time) remained, to assess motivation and reward. We acknowledge that questionnaires are limited in their ability to determine the underlying cause of behavioral change. For example, answering “he/she shows less enthusiasm for his or her usual interests” may be confounded by learned restrictions arising from physical motor impairments, or be influenced by semantic impairments and executive deficits. By using many tasks across a number of populations, we suggest that the extracted dimensions of apathy and impulsivity more accurately capture the essence of these behavioral changes than the use of single questions or tasks in isolation.

Finally, although PiPPIN aimed to be representative of the full population of affected patients, some may not have a diagnosis or be in contact with referring services. We also rely on clinical

Figure 2 White matter changes associated with carer-rated everyday skills (component 2) and carer-rated complex behaviors (component 3)
diagnostic criteria and acknowledge that some variants (svPPA, PSP) have much stronger clinic–pathologic correlations than others (CBS, bvFTD). Although we used multiple sources of referral in community and specialist services to reach all patients within the catchment area, some may have been missed. Nonetheless, the imaging subset was similar to the whole cohort.

White matter is markedly abnormal in the clinical syndromes associated with FTLD. DTI was sensitive to the white matter changes underlying FTLD-associated behaviors and revealed distinct spatial profiles relating to different aspects of apathy and impulsivity. These complex, multifaceted constructs are common across the FTLD spectrum and remain poorly treated. Elucidating the neural correlates of apathy and impulsivity, transdiagnostically, will help to inform the design of clinical trials for novel therapeutic strategies.

Author contributions
Conception and design of the study: J.B.R., I.T.S.-C.G., T.W.R. Acquisition of data: I.T.S.-C.G., C.J.L., J.B.R., P.V.R., E.W., K.M.D., P.S.J., A.W. Analysis of data: C.J.L., P.S.J. Drafting a significant portion of the manuscript: C.J.L., J.B.R.

Study funding
This work was funded by the NIHR Cambridge Biomedical Research Centre, the Cambridge Home and EU Scholarship Scheme, the James S. McDonnell Foundation (21st Century Science Initiative for Understanding Human Cognition), Wellcome Trust (103838), Medical Research Council (MC US A060 30PQ and RG62761), the Cambridge Brain Bank, PSP Association, and the Evelyn Trust. The BCNI is supported by a joint award from the Wellcome Trust and Medical Research Council. The authors thank the PSP Association and FTD Support Group for raising awareness of the study.

Disclosure
C. Lansdall, I. Coyle-Gilchrist, P. Jones, P. Vázquez Rodríguez, A. Wilcox, E. Wehmann, and K. Dick report no disclosures relevant to the manuscript. T. Robbins: consultancy for Cambridge Cognition, Lundbeck, Mundipharma, and Otsuka; research grants from Lundbeck and Shionogi; royalties for CANTAB from Cambridge Cognition; editorial honoraria for research grants from Lundbeck and Shionogi; royalties for Cambridge Cognition, Lundbeck, Mundipharma, and Otsuka; A. Wilcox, E. Wehmann, and K. Dick report no disclosures relevant to the manuscript. T. Robbins: consultancy for Cambridge Cognition, Lundbeck, Mundipharma, and Otsuka; research grants from Lundbeck and Shionogi; royalties for CANTAB from Cambridge Cognition; editorial honoraria from Psychopharmacology (Springer) and Current Opinion in Behavioral Sciences (Elsevier). J. Rowe: consultancy for Asce-neuron; research grants from AZ-MedImmune; serves as editor for Brain. Go to Neurology.org/N for full disclosures.

Received July 31, 2017. Accepted in final form December 21, 2017.

References
White matter change with apathy and impulsivity in frontotemporal lobar degeneration syndromes

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Cite as: Neurology® 2018;90:e1066-e1076. doi:10.1212/WNL.0000000000005175

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Study question
Do white matter (WM) abnormalities correlate with apathy and impulsivity in disorders associated with frontotemporal lobar degeneration (FTLD)?

Summary answer
Apathy and impulsivity are associated with distinct structural network changes in the WM of patients with FTLD-related syndromes.

What is known and what this paper adds
Apathy and impulsivity are common consequences of FTLD, and understanding their neurobiological basis may aid the development of neuroprotective strategies. This study clarifies the WM tract abnormalities associated with specific dimensions of apathy and impulsivity.

Participants and setting
This study examined 69 patients with all major syndromes of FTLD, drawn from a larger epidemiologic cohort, including behavioral variant frontotemporal dementia, primary progressive aphasia, progressive supranuclear palsy, and corticobasal syndrome, and 28 age- and sex-matched healthy controls.

Design, size, and duration
All participants underwent an extensive battery of neuropsychological and behavioral tests designed to assess various aspects of apathy and impulsivity. They also underwent diffusion tensor imaging (DTI) at 3 T. The DTI data were used for tract-based spatial analysis with threshold-free cluster enhancement (TFCE) corrections.

Primary outcomes
The primary outcomes were correlations between cognitive and behavioural profiles and DTI-detected WM tract abnormalities.

Main results and the role of chance
The study found that carer-rated changes in everyday skills, self care and apathy were associated with WM abnormalities in the corpus callosum, corona radiata, corticospinal tract, and posterior thalamic radiation (TFCE-corrected $p < 0.01$ for all). Complex behaviors including impulsivity correlated with abnormal frontotemporal connections between the orbital- and ventrolateral-prefrontal cortex, anterior cingulate, and temporal pole (TFCE-corrected $p < 0.01$ for all). Task-related impulsive behaviors correlated with changes in regions connecting the pre-supplementary motor area and dorsolateral prefrontal cortex, and the occipital lobe (TFCE-corrected $p < 0.05$ for all).

Bias, confounding, and other reasons for caution
DTI indirectly measures the physical properties of the brain parenchyma and is potentially subject to motion and registration errors. The neuropsychological battery was necessarily selective. Pathology is unknown in these patients.

Generalizability to other populations
The study population was drawn from a larger epidemiologic cohort of FTLD syndromes in the UK. Genetic and cultural variation between countries may constrain generalization of the results.

Study funding/potential competing interests
This study was funded by the British and EU governments and various medical research foundations. Prof. Robbins and Prof. Rowe report receiving consultancy fees and grants from various companies and having editorial relationships with various journals. Go to Neurology.org/N for full disclosures.

A draft of the short-form article was written by M. Dalefield, a writer with Editage, a division of Cactus Communications. The authors of the full-length article and the journal editors edited and approved the final version.
White matter change with apathy and impulsivity in frontotemporal lobar degeneration syndromes
Claire J. Lansdall, Ian T.S. Coyle-Gilchrist, P. Simon Jones, et al.
Neurology 2018;90:e1066-e1076 Published Online before print February 16, 2018
DOI 10.1212/WNL.0000000000005175

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