Terra incognita—cerebellar contributions to neuropsychiatric and cognitive dysfunction in behavioral variant frontotemporal dementia

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Although converging evidence has positioned the human cerebellum as an important relay for intact cognitive and neuropsychiatric processing, changes in this large structure remain mostly overlooked in behavioral variant frontotemporal dementia (bvFTD), a disease which is characterized by cognitive and neuropsychiatric deficits. The present study assessed whether degeneration in specific cerebellar subregions associate with indices of cognition and neuropsychiatric performance in bvFTD. Our results demonstrate a relationship between cognitive and neuropsychiatric decline across various domains of memory, language, emotion, executive, visuospatial function, and motivation and the degree of gray matter degeneration in cerebellar lobules V–VII. Most notably, bilateral cerebellar lobule VII and the posterior vermis emerged as distinct for memory processes, the right cerebellar hemisphere underpinned emotion, and the posterior vermis was highlighted in language dysfunction in bvFTD. Based on cortico-cerebellar connectivity maps, these findings in the cerebellum are consistent with the neural connections with the cortices involved in these domains in patients with bvFTD. Overall, the present study underscores the significance of cortical-cerebellar networks associated with cognition and neuropsychiatric dysfunction in bvFTD.

Keywords: cerebellum, behavioral variant frontotemporal dementia, cognition, neuropsychiatric processes, neural correlates

Introduction

The human cerebellum has long been regarded as an important relay station for intact motor function but converging evidence has now established its significant involvement also in cognitive and neuropsychiatric processes (Krienen and Buckner, 2009; O’Reilly et al., 2010; Stoodley and Schmahmann, 2010). It is perhaps of some surprise then that changes in this large brain region remain mostly overlooked in behavioral variant frontotemporal dementia (bvFTD), a disease characterized by neuropsychiatric and cognitive deficits, and that demonstrates motor features of amyotrophic lateral sclerosis (ALS) in a subpopulation of patients (Bak and Hodges, 2001). Regardless of the presence of concomitant ALS (Lillo et al., 2010), the progressive deterioration
in behavior and personality in bvFTD (Rascovksy et al., 2011) has been largely attributed to degeneration in the prefrontal, insular and temporal cortices (Rabinovici et al., 2007; Seeley et al., 2008). These cortical regions are known to demonstrate dense reciprocal connections with the cerebellum (Middleton and Strick, 2000, 2001; O’Reilly et al., 2010) and we recently confirmed that the degeneration of particular cerebellar subregions impacts on the overall cognitive and neuropsychiatric performances in bvFTD (Tan et al., 2014). However, the involvement of cerebellar substructures in specific cognitive and neuropsychiatric processes in bvFTD has yet to be examined.

The present study set out to extend on our previous findings of cerebellar contributions to global cognitive and neuropsychiatric measures, and examine the involvement of cerebellar subregions to more specific indices of cognition (memory, language, executive, emotion, visuospatial function) and neuropsychiatric performance (abnormal behavior, motivation, stereotypic behavior, mood, eating habits, and beliefs). A functional topographical map has been proposed in the human cerebellum, with lobules I–V and VIII found to be involved in sensorimotor function, contralateral lobules VI and VII in cognitive processing, and the posterior vermis conveying the limbic cerebellum (Krienen and Buckner, 2009; O’Reilly et al., 2010; Stoodley and Schmahmann, 2010; Stoodley et al., 2012). Based on these connectivity studies as well as the neural correlates of cognitive and neuropsychiatric domains identified in the cortices (Grossman et al., 2004; Huey et al., 2009; Raczka et al., 2010; Pennington et al., 2011; Kumfor et al., 2013; Woost et al., 2013; Irish et al., 2014) we predicted significant involvement of cerebellar lobules VI and VII across neuropsychiatric and cognitive domains with greater right lateralized involvement in executive function and language, and left lateralized involvement in visuospatial function in bvFTD.

Methods

Case Selection

A total of 53 participants took part in this study. Patients were sourced from the FTD Research Clinic, FRONTIER, resulting in a sample of 23 bvFTD, 15 ALSFTD, and 15 controls. All FTD patients met current consensus criteria for bvFTD (Rascovksky et al., 2011), showing the progressive behavioral and/or cognitive decline characteristic of this dementia subset, including some to all of the following: disinhibition, apathy, inertia, loss of empathy, perseveration, stereotypic behaviors and dysexecutive syndrome. Patients also met criteria of evidence of atrophy localized to anterior and/or temporal lobes via MRI. The ALSFTD group comprised patients who met diagnostic criteria for both bvFTD and ALS according with El Escorial Criteria Revised (Brooks et al., 2000), showing both upper and lower motor neuron signs and progressive behavioral/cognitive dysfunction. Well-matched healthy controls were recruited from the Frontier database. Testing and scanning was conducted at the first clinic visit of each patient. Only two bvFTD patients of the included cohorts had the C9orf72 mutation.

Cognitive and Neuropsychiatric Assessments

All participants were assessed on the Addenbrook's Cognitive Examination Revised (ACE-R) (Mioshi et al., 2006) as a measure of general cognitive ability. Specific cognitive assessments were performed as follows: The Rey Auditory Verbal Learning Test (RAVLT) was administered as an index of verbal recall and recognition, the Rey-Osterrieth Complex Figure Test (RCF) was used to assess visual recall, and the Doors and People test (part A) a measure of visual recognition. These three tasks have been described in greater detail previously (Pennington et al., 2011). The Boston Naming Test (BNT) (Goodglass et al., 2000) and Sydney Language Battery (SYDBAT) (Savage et al., 2013) were administered as standardized indices of verbal semantic performance. The Digit Span Reverse and FAS Verbal Fluency tests were used as measures of executive performance as described previously (Hornberger et al., 2008). Participants were given the Ekman 60 Faces Test obtained from the Facial Expressions of Emotion—Stimuli and Tests (FEEST) (Young Aw et al., 2002) and the facial emotion recognition task, which comprises the facial perceptual task (FPT), facial identity discrimination task (FIDT), facial affect discrimination task (FADT), and facial affect selection task (FAST) as previously described (Miller et al., 2012; Kumfor et al., 2014). Behavioral disturbances in the patients were assessed via the Cambridge Behavioral Inventory (CBI), which is an 81 item questionnaire that assesses cognitive, behavioral and affective symptoms as well as activities of daily living and evaluates various functional/behavioral domains using a 5 point rating scale (Wedderburn et al., 2008). The following scores of the CBI-R were evaluated and analyzed: abnormal behavior, motivation, stereotypic behavior, mood, eating habits, and beliefs. For all assessments, only data collected at the time of neuroimaging scans were included in the analyses.

Composite Scores

All neuropsychological test scores were converted into percentage of the control mean, before averaging to yield composite scores. Scores for the RAVLT, RCF, and Doors and People test (part A) were averaged to produce a memory composite. Scores for the BNT and SYDBAT were averaged to produce a language composite. Composite scores for executive function were derived from the Digit Span Reverse and FAS Verbal Fluency test scores, and scores for emotion derived from the Ekman 60 Faces Test and emotion selection task.

Statistics

Data were analyzed using IBM SPSS 20.0. A priori, variables were plotted and checked for normality of distribution by Kolmogorov-Smirnov tests. Parametric demographic data (age, education), cognitive (ACE-R) and neuropsychiatric (CBI-R)
data were compared across the three groups (bvFTD, ALS-FTD, and controls) via One-Way ANOVAs followed by Tukey HSD post-hoc tests. Variables revealing non-normal distributions were log transformed and the appropriate log values were used in the analyses, but Table 1 reports their original values to facilitate clinical interpretation. Variables showing non-parametric distribution after log transformation were analyzed via Chi-square (gender) and Kruskal-Wallis & Mann-Whitney U (disease duration) tests.

### Imaging Acquisition and Voxel-Based Morphometry (VBM) Analysis

Subjects were scanned using a 3T Philips MRI scanner. T1-weighted acquisition: coronal orientation, matrix 256 × 256 × 200, 161 mm³ in-plane resolution, slice thickness 1 mm, TE/TI = 2.6/5.8 ms.

Voxel-based morphometry (VBM) was conducted on the three dimensional T1-weighted scans, using the FLS-VBM toolbox in the FMRIB software library package (http://www.fmrib.ox.ac.uk/fsl/). The first step involved extracting the brain

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**Table 1 | Demographics, cognition, and neuropsychiatric measures in ALSFTD, bvFTD, and control groups.**

| Variable                        | ALSFTD         | bvFTD          | Controls        |
|---------------------------------|----------------|----------------|-----------------|
| **DEMOGRAPHICS AND GENERAL COGNITION** |                |                |                 |
| Age (y)                         | 63 ± 7.6       | 62 ± 10.1      | 63 ± 4.9        |
| Education (y)                   | 13 ± 3.6       | 12 ± 3.1       | 14 ± 1.7        |
| Gender (M/F)                    | 10/5           | 15/8           | 7/8             |
| Disease duration (y)            | 3 ± 2.3        | 4 ± 2.4        | N/A             |
| ACE-R (total score: 0–100)      | 30 ± 38**      | 74 ± 31**      | 98 ± 43         |
| **EXECUTIVE**                   |                |                |                 |
| Composite score                 | 30.2 ± 22.9**  | 52.1 ± 25.7**  | 100 ± 21.0      |
| DigitSpan backwards             | 2.9 ± 2.5**    | 4.4 ± 2.5**    | 8.6 ± 2.5       |
| Letter fluency                  | 13.1 ± 11.6**  | 21.8 ± 11.6**  | 45 ± 12.8       |
| **LANGUAGE**                    |                |                |                 |
| Composite score                 | 67.1 ± 25.9*   | 80.3 ± 27.2*   | 99.9 ± 3.5      |
| Boston naming                   | 9.5 ± 3.3*     | 11.9 ± 2.8*    | 14.5 ± 0.8      |
| Sydney battery                  |                |                |                 |
| Naming                          | 14.6 ± 17.6*   | 19.0 ± 6.5*    | 27.4 ± 1.8      |
| Comprehension                   | 21.6 ± 6.3*    | 24.0 ± 5.4**   | 29.5 ± 0.8      |
| Semantic                        | 19.6 ± 8.9*    | 21.2 ± 6.3*    | 28.4 ± 1.1      |
| **MEMORY**                      |                |                |                 |
| Composite score                 | 48.3 ± 32.0**  | 53.7 ± 25.7**  | 100 ± 16.5      |
| RCF 3 min                       | 7.0 ± 8.2      | 8.5 ± 6.8*     | 17.8 ± 4.8      |
| RAVLT A6                        | 4.1 ± 4.3*     | 4.0 ± 2.9**    | 9.6 ± 2.9       |
| Doors                           | 8.8 ± 2.7      | 7.7 ± 2.8*     | 11.4 ± 0.7      |
| **VISUOSPATIAL**                |                |                |                 |
| ACE-R (visuospatial: 0–100)     | 79 ± 17*       | 86 ± 15*       | 98 ± 4          |
| **EMOTION**                     |                |                |                 |
| Composite score                 | 71.7 ± 18.6**  | 73.2 ± 16.5**  | 100 ± 3.7       |
| Ekman60                         | 33.0 ± 14.2    | 35.6 ± 8.1     | 50.9 ± 4.5      |
| FPT                             | 30.7 ± 8.1     | 32.8 ± 8.9*    | 39.2 ± 1.4      |
| FIDT                            | 26.0 ± 5.0*    | 26.3 ± 6.4**   | 36.1 ± 4.3      |
| FAST                            | 22.2 ± 9.0*    | 26.9 ± 9.2**   | 39.0 ± 1.7      |
| FADT                            | 26.3 ± 5.5*    | 28.5 ± 6.8*    | 36.9 ± 1.5      |
| **CBI**                         |                |                |                 |
| Abnormal behavior (0–100)       | 25.3 ± 5.9*    | 36.1 ± 4.8**   | 1.1 ± 6.6       |
| Motivation (0–100)              | 36.1 ± 8.3**   | 66.7 ± 6.8**   | 2.3 ± 9.3       |
| Stereotypic behavior (0–100)    | 45.1 ± 6.8*    | 54.2 ± 5.6**   | 6.3 ± 7.7       |
| Mood (0–100)                    | 21.0 ± 5.3     | 34.2 ± 4.4**   | 5.5 ± 6.0       |
| Eating (0–100)                  | 29.5 ± 5.9     | 47.6 ± 4.8**   | 7.4 ± 6.7       |
| Everyday skills (0–100)         | 19.3 ± 5.8     | 30.5 ± 4.7*    | 0.5 ± 6.5       |
| Beliefs (0–100)                 | 1.8 ± 4.1      | 9.9 ± 3.3      | 0.0 ± 4.6       |

Data are presented as mean ± standard deviation. Composite scores are presented as a % of mean control. Differences between groups are represented as *p < 0.05 compared to controls; **p < 0.01 compared to controls; ***p < 0.05 compared to bvFTD; FPT, facial perceptual task; FIDT, facial identity discrimination task; FADT, facial affect discrimination task; FAST, facial affect selection task.

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**Figure 1 | Voxel-based morphometry analyses showing cerebellar regions in which gray matter intensity correlates significantly with memory, language, executive, emotion, visuospatial task performance, and motivation measures across all participant groups.** Colored voxels show regions that were significant in the analyses for p < 0.01 uncorrected and a cluster threshold of 20 contiguous voxels. All clusters reported t > 3.5. Clusters are overlaid on the MNI standard brain with a mask for lobule VII (crus 1, 2, and VIIb) shown in blue and a mask for the vermis shown in light blue. L, Left Hemisphere; R, Right Hemisphere.
from all scans using the BET algorithm in FSL, using a fractional intensity threshold of 0.22 (Smith, 2002). Each scan was visually checked after brain extraction, both to ensure that no brain matter was excluded, and no non-brain matter was included (e.g., skull, optic nerve, dura mater).

If non-brain matter was visually detected or brain matter was falsely excluded, the BET algorithm for that scan was repeated with a modified fractional intensity threshold, to give smaller or larger brain border estimates. A gray matter template, specific to this study, was then built from canvassing 10 scans from each group (total \( n = 30 \)). An equal amount of scans across groups was used to ensure equal representation, and thus avoid potential bias toward any single group's topography during registration. Template scans were then registered to the Montreal Neurological Institute Standard space (MNI 152) using non-linear b-spline representation of the registration warp field, resulting in study-specific gray matter template at \( 2 \times 2 \times 2 \text{mm}^3 \) resolution in standard space. Simultaneously, brain-extracted scans were also processed with the FMRIB’s Automatic Segmentation Tool (FAST v4.0) (Zhang et al., 2001) to achieve tissue segmentation into CSF, gray matter, and white matter. Specifically this was done via a hidden Markov random field model and an associated Expectation-Maximization algorithm. The FAST algorithm also corrected for spatial intensity variations such as bias field or radio-frequency inhomogeneities in the scans, resulting in partial volume maps of the scans. The following step saw gray matter partial volume maps then non-linearly registered to the study-specific template via non-linear b-spline representation of the registration warp. These maps were then modulated by dividing by the Jacobian of the warp field, to correct for any contraction/enlargement caused by the non-linear component of the transformation (Andersson et al., 2007a). After normalization and modulation, smoothing the gray matter maps occurred using an isotropic Gaussian kernel (standard deviation = 3 mm; full width half maximum = 8 mm). The statistical analysis was performed with a voxel-wise general linear model after correcting for intracranial volume. Significant clusters were formed by employing the threshold-free cluster enhancement (TFCE) method (Smith and Nichols, 2009). The TFCE method is a cluster-based thresholding method which does not require the setting of an arbitrary cluster forming threshold (e.g., \( t, 2 \)). Instead, it takes a raw statistics image and produces

### TABLE 2 | Voxel-based morphometry (VBM) findings demonstrating gray matter volumes in cerebellum subregions showing a significant correlation with measures of memory, language, executive, emotion, visuospatial task performances (\( p < 0.01 \)) shown in Figure 1.

| Cerebellar subregions | Memory | Language | Executive | Emotion | Visuospatial |
|-----------------------|--------|----------|-----------|---------|--------------|
| I–IV                  |        |          |           |         |              |
| V                     |        |          |           |         |              |
| VI                    |        |          |           |         |              |
| VII (Crus 1)          |        |          |           |         |              |
| VII (Crus 2)          |        |          |           |         |              |
| VIIIB                 |        |          |           |         |              |
| VIII                  |        |          |           |         |              |
| IX–X                  |        |          |           |         |              |
| Vermis                |        |          |           |         |              |

Solid gray color represents areas of substantial correlation, diagonal lines represent areas of milder significance.

### TABLE 3 | Voxel-based morphometry (VBM) findings demonstrating shared regions of cerebellar gray matter atrophy for memory (brown color), language (blue color), emotion (green color), executive (pink color), and visuospatial performances (red color) across all participants (\( p < 0.01 \)).

| Cerebellar subregions | Memory | Language | Executive | Emotion | Visuospatial |
|-----------------------|--------|----------|-----------|---------|--------------|
| I–IV                  | R L R L | R L R L  | R L R L   | R L R L  |              |
| V                     | R L R L | R L R L  | R L R L   | R L R L  |              |
| VI                    | R L R L | R L L L  | R L L L   | R L R L   | L L L L     |
| VII (Crus 1)          | R L R L | R L L L  | R L R L   | R L R L   | L L L L     |
| VII (Crus 2)          | R L R L | R L L L  | R L R L   | R L R L   | L L L L     |
| VIIIB                 | R L R L | R L L L  | R L R L   | R L R L   |              |
| VIII                  | R L    |          |           |         |              |
| IX–X                  |        |          |           |         |              |
| Vermis                | R L    |          |           |         |              |

| Memory | Language | Executive | Emotion | Visuospatial |
|--------|----------|-----------|---------|--------------|
| Memory |          | R L       |         |              |
| Language |        |          | R L     |              |
| Executive |       |          |         | R L         |
| Emotion |          |          |         |             |
| Visuospatial | |          |         |              |
an output image in which the voxel-wise values represent the amount of cluster-like local spatial support. The TFCE image is then turned into voxel-wise $p$-values via permutation testing. We employed a permutation-based non-parametric testing with 5000 permutations (Nichols and Holmes, 2002).

Region-of-Interest Mask
A region-of-interest (ROI) mask was created for subregions of the human cerebellum using the Schmahmann probabilistic atlas of the human cerebellum (http://www.icn.ucl.ac.uk/motorcontrol/imaging/propatlas.htm) as previously described (Tan et al., 2014).

VBM: Correlations with Test Scores
In a first step, correlations between performance on cognitive and neuropsychiatric assessments with regions of gray matter atrophy in the cerebellum were investigated across participant groups by entering the total scores of ACE-R and CBI-R, and the composite scores of language, memory, emotion, executive and visuospatial assessments as a covariate in the design matrix of the VBM analysis of the cerebellum across participant groups. For statistical power, a covariate only statistical model with a (1) t-contrast was used, providing an index of association between gray matter intensity and clinical scores. In a second analysis, we performed an inclusive masking analysis to verify which areas of cerebellar atrophy that correlate with language, memory, emotion, executive, and visuospatial scores would overlap across cognitive domains. In a final step, we performed an exclusive masking analysis to determine which areas of cerebellar atrophy were specific to language, memory, emotion, executive, and visuospatial scores.

Anatomical locations of significant results were overlaid on the MNI standard cerebellum image for spatial normalization and visual comparison with a brain atlas, allowing localization of areas of significant gray matter loss. The covariate analyses were tested for significance at $p < 0.01$ uncorrected with a voxel cluster threshold of 20 contiguous voxels. Given that cerebellar volumes in these same patient cohorts had already been contrasted previously Supplementary Figure 1 (Tan et al., 2014), group differences were not assessed in the present study.

Results

Demographics
Demographic and clinical information is presented in Table 1. The participant groups were well-matched for age ($p = 0.66$), years in education ($p = 0.32$), gender ($p = 0.78$), and disease duration (i.e., the number of months elapsed between onset of symptoms and cognitive testing, $p = 0.88$).

Test Scores
Table 1 details cognitive and neuropsychiatric assessments as well as composite scores for each task category across participant groups. Across the various cognitive and neuropsychiatric assessments administered, a significant poorer performance was seen in both patient groups in comparison to controls (Table 1). No significant difference was identified between patient cohorts with the exception of the motivation component of the CBI-R assessment, where a poorer performance was seen in patients with bvFTD ($p < 0.05$).

Neural Correlates of Cognitive and Neuropsychiatric Scores
As can be seen in Figure 1 and Table 2, correlations between cerebellar gray matter atrophy with cognitive and neuropsychiatric measures were examined across all groups. Memory scores (brown color, Figure 1 and Table 2) correlated significantly with gray matter volumes in the posterior lobules VI–VIIB and vermis, with mild involvement identified in anterior lobules I–V. Composite scores of language (blue color, Figure 1 and Table 2) correlated significantly with gray matter volumes in the posterior lobule VI, left lobule VII (Crus I) and the vermis, with mild involvement also seen in lobule V.
Executive measures (pink color, Figure 1 and Table 2) were found to associate with gray matter volumes in the posterior lobule VI and VII (Crus I), with a mild association seen in lobule V. Emotion scores (green color, Figure 1 and Table 2) correlated with gray matter atrophy in the posterior lobule VI and Crus I. Visuospatial measures (red color, Figure 1 and Table 2) correlated with gray matter volume in the left cerebellar lobule VI. Patients’ performance on tasks of motivation (red-yellow color, Figure 1 and Table 2) showed a significant association with gray matter volume in right-lateralized lobule VI. No significant correlations were identified between cerebellar volumes with neuropsychiatric measures of abnormal behavior, motivation, stereotypic behavior, mood, eating habits, and beliefs.

Overlap Analysis
Regional gray matter in the cerebellum associated with memory, language, executive, emotion, and visuospatial scores were further investigated to determine regions of overlap with other cognitive domains. Table 3 and Figure 2 demonstrates significant overlap across all cognitive domains in the left cerebellar lobule VI, with further shared representations seen in the right cerebellar lobule VI and bilateral lobules V and VII (Crus I) for memory, language, executive, and emotion processes. A further shared representation was seen in the vermis shared between memory and language domains.

Difference Analysis
An exclusive analysis was performed to determine cerebellar regions specific to memory, language, executive, emotion, and visuospatial scores. Bilateral lobules VII (Crus I–II) and vermis emerged as distinct to memory performance (brown color, Table 4 and Figure 3); the posterior vermis was found to be exclusively involved in language dysfunction (blue color, Table 4 and Figure 3); mild involvement of right lobules VII (Crus I) and left lobule VI was exclusive to executive functioning (pink color, Table 4 and Figure 3); bilateral lobules VI–VII (Crus I) with greater right hemisphere involvement was exclusive to emotion processing (green color, Table 4 and Figure 3), and left lobule VI emerged as exclusive to visuospatial function (red color, Table 4 and Figure 3).

Discussion
The present study investigated the neural correlates of cognitive and neuropsychiatric performance in the cerebellum of patients with bvFTD. Our results show that cognitive and neuropsychiatric decline across various domains of memory, language, emotion, executive, visuospatial function, and motivation relates to the degree of gray matter degeneration in cerebellar lobules V–VII. Comparisons across task categories were performed to determine overlapping and distinct cerebellar subregions associated with deficits in each cognitive domain. This revealed a convergence across all cognitive and neuropsychiatric substrates on cerebellar lobule VI, with additional associations seen in lobules V and VII (Crus I) for memory, language, executive, and emotion deficits, and a further shared representation present between memory and language dysfunction in the vermis. Most notably, bilateral cerebellar lobules VII and vermis emerged as distinct for deficits in memory processing, the right cerebellar hemisphere associated with emotion dysfunction and the cerebellar vermis was highlighted in language dysfunction.

Although the cerebellum is now known to play a critical role in cognition and neuropsychiatric processes (Schmahmann and Sherman, 1998; Stoodley and Schmahmann, 2010), evidence of its contribution to the cognitive and neuropsychiatric symptoms in bvFTD is only beginning to surface (Tan et al., 2014). The present study demonstrated that degeneration in cerebellar lobules VI–VII impacts on memory, emotion, executive, and visuospatial performances in bvFTD, corroborating findings in healthy humans (Stoodley and Schmahmann, 2009; Keren-Happuch...
Contrary to the expectation that the vermis is less connected to language processes in healthy humans and while variable results with regards to whether language function maps predominantly onto the right, left or bilateral cerebellar hemispheres have been found (Krienen and Buckner, 2009; Stoodley and Schmahmann, 2010; Stoodley et al., 2012), we demonstrate here an association between bilateral lobule VI and left Crus I with language dysfunction in bvFTD. We further demonstrated involvement of the posterior vermis, also known as the “limbic cerebellum,” in deficits in the language and memory domains in bvFTD.

Given that intact cognition is multifaceted and contingent upon shared memory, language, executive, emotion, and visuospatial processes, cognitive substrates that each individually also draw upon another, we conducted comparisons across task categories to further identify regions significant to deficits in each domain. Overlap analyses was first performed to determine if within cerebellar subregions implicated across task categories, there existed regions of overlap. Our findings revealed collective clusters within lobules VI and Crus I across deficits in the memory, language, executive, emotion and visuospatial domains, and in the vermis across memory and language dysfunction. We then proceeded to perform comparisons to determine subregions distinct to each category within these cerebellar substructures. Within each of these individual domains, discriminate analysis underscored the bilateral lobules VII and vermis to memory performance, the vermis to language performance, right-lateralised lobules VI and VII (Crus I) to deficits in emotion, and left-lateralized lobule VI to visuospatial dysfunction. Only mild involvement of cerebellar lobules VI and VII (Crus I) was observed with executive dysfunction in the absence of memory, language, emotion and visuospatial substrates. While this may be due to the recruitment of other cognitive domains in the present assessments of executive dysfunction, this is not an isolated report of difficulties in identifying cerebellar regions involved purely in executive function (Stoodley and Schmahmann, 2009).

Intact memory in bvFTD has been found to be contingent upon the integrity of the prefrontal cortices and the limbic “Papez” cortical-subcortical network (Hornberger et al., 2012; Frisch et al., 2013; Irish et al., 2013). Given that connectivity studies have shown closed-loop connections between the Crus with the prefrontal cortices and the vermis with the limbic network (Middleton and Strick, 2001; Kelly and Strick, 2003; O’Reilly et al., 2010), the present findings of distinct lobule VII (Crus) and vermis contributions to memory performance in bvFTD underscores the significance of the cortical-cerebellar network to intact memory processing in bvFTD. Based on the left-hemispheric involvement of the cortical regions in language in bvFTD, we anticipated greater contributions from the right cerebellar hemisphere. However, our results revealed that in the absence of other cognitive task categories, only the vermis emerged as distinct to deficits in language processing in bvFTD, a finding that although not reported in healthy humans (Stoodley and Schmahmann, 2009), is observed in individuals with autism (Riva et al., 2013). Contrary to the expectation that the vermis would be more involved in deficits in emotion processing (Baumann and Mattingley, 2012), predominant right-lateralized lobule VI emerged as distinct with measures of emotion processing, a finding which may relate to the predominant left hemispheric recruitment of various cortices during these tasks in bvFTD (Kumfor et al., 2013; Bertoux et al., 2014).

Apathy is a prominent feature of bvFTD and studies performed so far suggest that the neural basis underlying this lack of motivation is the disruption of the cortical-basal ganglia circuitry in patients with bvFTD (Eslinger et al., 2012; Yi et al., 2013). Despite being a recipient and reciprocal relay of cortical projections via the basal ganglia, cerebellar involvement was not highlighted in these studies. However, one case report demonstrated a simultaneous reduction of apathy and improved metabolism in the bilateral insula and cerebellum with the administration of an N-methyl-D-aspartate antagonist to a patient with bvFTD (Links et al., 2013). Together with the implications of the current study for focal cerebellar atrophy in bvFTD, these findings underscore the need for further studies of apathy and cerebellar deficits in bvFTD.
present findings pinpointing right-lateralized cerebellar lobule VI in motivation, it appears that the cerebellum remains largely overlooked in the study of neuropsychiatric deficits in bvFTD, particularly in sporadic disease. There are several methodological issues that warrant consideration. In order to explore the cerebellar correlates of cognitive and neuropsychiatric measures in this group, covariate analyses was assessed for significance at \( p < 0.01 \) uncorrected and it was beyond the scope of this study to determine if the cerebellar correlates identified here are specific to bvFTD, or seen across other neurodegenerative diseases as well. However, despite this, it is important to note that our findings here converge well with a growing number of studies that have highlighted cerebellar subregions involved in cognitive and neuropsychiatric measures across cohorts of healthy humans and other lesion models. Finally, cerebellar correlations were not assessed across each bvFTD cohort with controls due to (1) our findings both here (Table 1) and previously (Lillo et al., 2010) of no significant difference in cognitive and neuropsychiatric measures across these two bvFTD cohorts; (2) the substantial body of literature demonstrating that bvFTD and ALSFTD sit on the same disease continuum; (3) sample size \( n = 23 \) bvFTD, \( n = 15 \) ALSFTD) teamed with the above two rationale and (4) the focus being cerebellar atrophy with cognitive and neuropsychiatric changes rather than motor impairment which, in the case of the latter, would warrant analyses to have been performed across each patient group with controls. However, it would be of interest to perform covariation analyses within larger participant cohorts in the future.

In summary, the present findings of significant cerebellar gray matter associations with cognitive and neuropsychiatric processes in bvFTD are consistent with the neural correlates identified with these domains in the cortical regions involved. This emphasizes the involvement of whole brain cortical-cerebellar network in cognition and neuropsychiatric function, highlighting the importance of including cerebellar assessment in whole brain analyses of structural changes in bvFTD.

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Supplementary Material

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fnagi.2015.00121

Supplementary Figure 1 | Voxel-based morphometry findings contrasting gray matter atrophy in bvFTD and ALSFTD in comparison to controls, and bvFTD in comparison to ALSFTD. Colored voxels show regions that were significant in the analysis for \( p < 0.05 \) family-wise error (FEW) corrected. Results are published in PLoS ONE 9(8): e105632. doi:10.1371/journal.pone.0105632 and demonstrate significant cerebellar atrophy in both bvFTD and ALSFTD cohorts in comparison to controls, but no significant difference across patient groups. Clusters are overlaid on the MNI standard brain with a mask for lobule VII (crus 1, 2, and Vb) shown in blue.

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