ABSTRACT

Objective: To identify distinct behavioral phenotypes of behavioral-variant frontotemporal dementia (bvFTD) and to elucidate differences in functional, neuroimaging, and progression to residential care placement.

Methods: Eighty-eight patients with bvFTD were included in a cluster analysis applying levels of disinhibition and apathy (Cambridge Behavioural Inventory-Revised) to identify phenotypic subgroups. Between-group (Kruskal-Wallis, Mann-Whitney U) functional differences (Disability Assessment for Dementia) and time to residential care placement (survival analyses) were examined. Cortical thickness differences (whole-brain MRI) were analyzed in patients with bvFTD vs healthy controls (n = 30) and between phenotypic subgroups.

Results: Four phenotypic subgroups were identified: primary severe apathy (n = 26), severe apathy and disinhibition (n = 26), mild apathy and disinhibition (n = 27), and primary severe disinhibition (n = 9). Patients with severely apathetic phenotypes were more functionally impaired and had more extensive brain atrophy than those with mild apathy or severe disinhibition alone. Further imaging analyses indicated that the right middle temporal region is critical for the development of disinhibition, an association that remains with disease progression and in the context of severe apathy. Finally, no difference in time to residential care admission was found between phenotypes.

Conclusions: This study reveals that different clinical behavioral phenotypes of bvFTD have differing profiles of functional decline and distinct patterns of associated cortical changes. These findings emphasize the importance of apathy in functional impairment, highlight the role of the right temporal region in disinhibition, and suggest that disability may be a sensitive outcome measure for treatments targeting reduction of apathy. These phenotypes could also support understanding of prognosis and clinical management. Neurology® 2017;89:1–8

GLOSSARY

ACE-III = Addenbrooke’s Cognitive Examination-III; ACE-R = Addenbrooke’s Cognitive Examination-Revised; ADL = activity of daily living; bvFTD = behavioral-variant frontotemporal dementia; CBI-R = Cambridge Behavioural Inventory-Revised; CI = confidence interval; DAD = Disability Assessment for Dementia; MA + D = mild apathy and disinhibition; PSA = primary severe apathy; PSD = primary severe disinhibition; ROI = region of interest; SA + D = severe apathy and disinhibition.

Behavioral symptoms are salient in behavioral-variant frontotemporal dementia (bvFTD), yet considerable phenotypic variability exists within the diagnosis.¹ Clinical studies have paid particular attention to the symptoms of apathy and disinhibition, which have been identified as causing the greatest caregiver distress.²³ bvFTD has been classified into apathetic (primary symptoms of aspontaneity, inertia, and slowness) or disinhibited (more distractibility, overactivity, and restlessness) behavioral presentations.⁴ Subsequent studies investigated associations of these behavioral phenotypes with metabolic and neuroanatomic characteristics.²³⁵–¹⁰

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Cognitively, apathetic bvFTD phenotypes have frequently been reported as more impaired than disinhibited,7,11 but there is no consensus at present.12 The functional implication of behavioral phenotypes on activities of daily living (ADLs) and prognosis has not been fully understood. Relative preservation of instrumental ADLs across both phenotypes has been reported,11 but instrumental ADLs have been subsequently described as more impaired in apathetic phenotypes.7,13 Indeed, increasing apathy has been correlated with declining functional ability in bvFTD.13,14

This study aimed to identify different behavioral profiles within bvFTD in a data-driven approach. We hypothesized that apathy, but not disinhibition, would be a strong contributor to functional disability; that severe apathy would be associated with a greater spread of cortical atrophy; and that severe apathy would contribute to more rapid placement to residential care.

METHODS Participants. Eighty-eight individuals diagnosed with bvFTD consecutively recruited by FRONTIER, the Fronto-temporal Dementia Research Group in Sydney (Australia), were included in the study. These individuals were compared with 30 age-, sex-, and education-matched healthy controls. Patients were assessed in clinic between November 2007 and June 2015. Patients were included if they met current criteria for either possible or probable bvFTD7 (diagnoses were based on a multidisciplinary consensus, neurologist and neuropsychologist), had a reliable proxy informant to report on their behavior and everyday routine, did not have major depression or other neuropsychiatric disease, and did not have physical limitations that could affect ADLs. Disease duration was estimated at the time of diagnosis from the onset of symptoms as described by the caregiver. Controls were recruited from the Neuroscience Research Australia Volunteer database. Healthy controls scored above 88 of 100 on the Addenbrooke Cognitive Examination-III (ACE-III)15 and 0 on the Sum of Boxes subscales (disinhibition and apathy from the CBI-R) to determine the optimal number of clusters for the data.21 Analysis of the change in distance between the cluster mergers on the dendrogram generated from this analysis (figure e-1) suggested that 4 clusters were an appropriate solution for this data.22 This 4-cluster solution was validated with 2 consecutive 2-step clustering procedures using euclidean distance, each with a different information criterion (appendix e-1 gives details of the procedure). The 4 clusters were then used in the next phase of analyses to compare the bvFTD subgroups (clusters) in terms of their ADL functioning.

Demographic data across the 4 cluster groups were compared with Kruskal-Wallis tests and χ2 tests for sex comparisons. Kruskal-Wallis tests were also used to explore differences across clusters for each DAD area (total DAD, basic ADLs, instrumental ADLs, initiation, planning, execution). Mann-Whitney U tests were then conducted for post hoc analyses with the Benjamini-Hochberg procedure used for false discovery rate control of multiple comparisons.23

Finally, to understand if belonging to different bvFTD subgroup clusters would lead to more rapid placement to residential care, a Kaplan-Meier survival analysis was conducted. Placement to residential care (survival time) was defined as time from symptom onset to placement into residential care. Censoring was used for patients who were still living at home at the time of analysis or who remained at home until their death. All statistical analyses were performed with SPSS 21.0 (IBM, Armonk, NY).

Imaging data analyses. Whole-brain structural MRIs were acquired and processed for bvFTD (n = 72) and healthy controls (n = 30) as described in appendix e-1. Four sets of vertex-by-vertex analyses were performed with general linear models to examine whole-brain differences in cortical thickness between the bvFTD subgroups resulting from the cluster analysis and healthy controls. Statistical significance was set at p = 0.001 uncorrected for multiple comparisons. Furthermore, direct comparisons between bvFTD subgroups were carried out with t test analyses to isolate the neural correlates of disease severity and disinhibition associated with the different clinical phenotypes. Statistical significance was set at p = 0.01 uncorrected for multiple comparisons in these pairwise comparisons. A conservative cluster extent threshold of k > 50 mm2 was used to minimize type I error while balancing the risk of type II error.24

Next, 4 greater cortical regions of interest (ROIs), one for each of the cortical lobes (frontal, temporal, parietal, and occipital), were selected, computed, and averaged across both hemispheres (for details, see table e-1). One-way analysis of variance was used to explore between-group differences in the average cortical thickness within these ROIs. Post hoc analyses were corrected for multiple comparisons with a Sidak adjustment. Finally, regression analyses were used to test for significant linear effects within these ROIs across the different groups. The level of significance was set at p < 0.05.

RESULTS bvFTD phenotypes: Cluster demographics. Clusters and healthy controls were matched for age, sex distribution, and education (table 1). All cluster groups were matched for disease duration; healthy controls had better cognitive scores overall.

Are there distinct bvFTD phenotypes? Cluster solution results. Figure 1 shows the 4 patient subgroups, which were labeled to reflect clinical symptoms. Primary severe apathy (PSA; n = 26) was characterized by...
severe apathy scores and mild disinhibition scores; severe apathy and disinhibition (SA + D; n = 26) was characterized by severe apathy and severe disinhibition scores; mild apathy and disinhibition (MA + D; n = 27) was characterized by mild apathy and mild disinhibition; and primary severe disinhibition (PSD; n = 9) was characterized by mild apathy and severe disinhibition. Group differences confirmed the distinction of these 4 subgroups for both apathy [χ²(3) = 64.641, p < 0.001] and disinhibition [χ²(3) = 63.122, p < 0.001] CBI-R scores. Post hoc tests revealed specific cluster group differences (figure 1).

**Functional disability differences across bvFTD phenotypes.** Group differences were identified for every area of daily function measured: overall ADLs [χ²(3) = 14.753, p < 0.005], basic ADLs [χ²(3) = 11.719, p < 0.01], instrumental ADLs [χ²(3) = 13.236, p < 0.005], and the 3 subcomponents of activity performance: initiation [χ²(3) = 16.068, p < 0.005], planning [χ²(3) = 9.957, p < 0.05], and execution [χ²(3) = 10.391, p < 0.05]. Post hoc analyses compared each cluster against the MA + D cluster (the least impaired phenotype). Next, the PSA and PSD clusters were compared. After multiple comparisons were controlled for, the PSA and the SA + D subgroups were more functionally impaired than the MA + D group for every functional area. No differences between the PSA and SA + D subgroups were found. Clinically, this translates as people who had severe

### Table 1  Demographics for bvFTD behavioral phenotypes (cluster groups) and healthy controls

<table>
<thead>
<tr>
<th></th>
<th>bvFTD all (n = 88)</th>
<th>PSA (n = 26)</th>
<th>SA + D (n = 26)</th>
<th>MA + D (n = 27)</th>
<th>PSD (n = 9)</th>
<th>Healthy controls (n = 30)</th>
<th>Across clusters and healthy controls</th>
<th>Across clusters only</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y</td>
<td>62.15 (8.68)</td>
<td>63.96 (7.93)</td>
<td>60.00 (10.52)</td>
<td>62.30 (7.28)</td>
<td>62.67 (8.87)</td>
<td>64.53 (4.19)</td>
<td>5.30 0.258a</td>
<td>2.52 0.471a</td>
</tr>
<tr>
<td>Sex (M/F), n</td>
<td>54/34</td>
<td>15/11</td>
<td>16/10</td>
<td>17/10</td>
<td>6/3</td>
<td>12/18</td>
<td>4.42 0.353b</td>
<td>0.28 0.963b</td>
</tr>
<tr>
<td>Education, y</td>
<td>11.87 (2.96)</td>
<td>11.82 (3.31)</td>
<td>11.44 (2.58)</td>
<td>12.28 (3.00)</td>
<td>12.00 (3.16)</td>
<td>12.35 (1.22)</td>
<td>5.29 0.259a</td>
<td>0.81 0.846a</td>
</tr>
<tr>
<td>Disease duration, y</td>
<td>4.14 (2.42)</td>
<td>4.47 (2.40)</td>
<td>4.62 (2.63)</td>
<td>3.45 (2.37)</td>
<td>3.99 (1.82)</td>
<td>NA</td>
<td>NA NA NA</td>
<td>4.31 0.230a</td>
</tr>
<tr>
<td>Baseline cognitive assessment; maximum 100, cutoff 88/100</td>
<td>65.54 (14.83)</td>
<td>61.22 (14.07)</td>
<td>62.59 (18.25)</td>
<td>70.41 (9.74)</td>
<td>71.50 (15.00)</td>
<td>95.23 (3.58)</td>
<td>64.77 &lt;0.001a</td>
<td>6.88 0.076a</td>
</tr>
</tbody>
</table>

Abbreviations: bvFTD = behavioral-variant frontotemporal dementia; MA + D = mild apathy and disinhibition; SA = primary severe apathy; PSD = primary severe disinhibition; SA + D = severe apathy and disinhibition.

Scores are means (SDs). Disease duration refers to the time between symptom onset and baseline assessment date.

*a* Kruskal-Wallis test E.

*b* Chi-square test.

### Figure 1  Levels of apathy and disinhibition across patient subgroups (clusters) identified in 2-step cluster analysis (with the Akaike criterion)

Higher scores indicate worse levels of behavioral expression. (A) PSA > MA + D and PSD (both U = 0.0, p < 0.001), SA + D > MA + D (U = 0.50, p < 0.001) and PSD (U = 0.0, p < 0.001). (B) SA + D > PSA and MA + D (both U = 0.0, p < 0.001), PSD > PSA (U = 3.50, p < 0.001) and MA + D (U = 0.50, p < 0.001). CBI = Cambridge Behavioural Inventory-Revised; MA + D = mild apathy and disinhibition; PSA = primary severe apathy; PSD = primary severe disinhibition; SA + D = severe apathy and disinhibition. Mann-Whitney U tests (Bonferroni correction); *p < 0.001.
Apathy were more disabled and had more difficulty with their everyday living tasks than those with mild apathy.

On the other hand, there were no differences between the PSD and MA D groups for any of the functional domains, indicating that, in contrast to those with a primarily apathetic presentation, patients with a primarily disinhibited presentation may retain some functional abilities. Surprisingly, there were no differences between the PSA and the PSD groups. Notably, the outlier included in the PSD group (n = 9) had a disease duration that was 5.26 years longer than the average disease duration for that group, had the equal highest concurrent levels of apathy in the PSD group, and scored zero in overall ADLs compared to the group mean of 60.57% (figure 2).

Imaging analyses. Group comparisons between bvFTD clusters and healthy controls revealed the characteristic profile of brain atrophy consistent with a diagnosis of bvFTD (figure e-2). In brief, all phenotypes (clusters) showed atrophy in the insula, inferior frontal, and anterior temporal cortices with some differences in the extent and location of the atrophy between subgroups. Patients in the SA + D and PSA clusters (marked apathy) exhibited an extensive and similar pattern of cortical thinning, with atrophy extending posteriorly to the temporoparietal junction and regions of the posterior cingulate cortex and precuneus. Patients in the MA + D cluster showed a similar pattern but with less extensive thinning. In contrast, patients in the PSD cluster (marked disinhibition) showed focal atrophy in the bilateral orbitofrontal, insular, and right anterior temporal and the anterior cingulate cortex.

Pairwise comparisons between the SA + D and PSA groups (both phenotypes, n = 26) were carried out to examine neural correlates of disinhibition, and SA + D and MA + D were compared to examine the neural correlates of apathy severity (figure 3). Disinhibition was associated with clusters of cortical thinning in the left temporal pole and right inferior and middle temporal gyri. Disease severity was associated with clusters of thinning in the left inferior temporal cortex and in the right insula, right middle temporal, and parietal cortices and right posterior cingulate/isthmus.

Finally, ROI analyses showed a linear trend between symptom load/severity and cortical thinning across the different clusters (from more to less severe cortical atrophy: SA + D, PSA, MA + D, and PSD; figure 4). These analyses yielded linear associations in the frontal ($R^2 = 0.25, p < 0.0001$), temporal ($R^2 = 0.19, p < 0.0001$), parietal ($R^2 = 0.10, p = 0.0014$), and occipital ($R^2 = 0.10, p = 0.0014$) lobes. Post hoc comparisons of mean cortical thickness revealed differences between patients from all clusters and healthy controls in the frontal and temporal lobes (both $p < 0.001$). In the parietal and occipital lobes, differences were observed between patients from the SA + D cluster and healthy controls (both $p < 0.05$). No differences were observed between patient subgroups.
bvFTD cluster phenotype and admission to residential care. At the time of the study, 38 participants had moved into residential care. One participant was excluded from the analysis because symptom onset information was unavailable. Therefore, 87 participants were included in the Kaplan-Meier survival analysis to compare the 4 patient groups in regard to admission to residential care. Overall, the median time in years from diagnosis to residential care admission for all the groups combined (n = 87) was 2.8 years (95% confidence interval [CI] 2.3–3.2). The median time from diagnosis to residential care admission for each group ranged between 1.3 and 3.6 years (PSA: 1.3 years, 95% CI 0.0–3.8; SA + D: 3.0 years, 95% CI 1.4–4.5; MA + D: 3.1 years, 95% CI 2.2–3.9; and PSD: 3.6 years, 95% CI 0.4–6.9), but there were no differences between groups (figure e-3; \( X_{(3)}^2 = 0.511, p = 0.916 \)).

Statistical significance was set at \( p = 0.01 \) uncorrected for multiple comparisons. bvFTD = behavioral-variant frontotemporal dementia; MA + D = mild apathy and disinhibition; PSA = primary severe apathy; SA + D = severe apathy and disinhibition.

Mean cortical thickness was averaged for both hemispheres. HC = healthy controls; MA + D = mild apathy and disinhibition; PSA = primary severe apathy; PSD = primary severe disinhibition; SA + D = severe apathy and disinhibition. \(* p < 0.05; ** p < 0.001\).
DISCUSSION This study confirms the heterogeneity of the bvFTD syndrome on the basis of severity of apathy and disinhibited symptoms. Moreover, these different bvFTD phenotypes have been translated into distinct everyday living functional profiles and are explained by different patterns of brain atrophy. We confirmed our hypothesis that apathy, but not disinhibition, had a greater negative effect on daily functioning. Despite clear functional differences between these behavioral phenotypes, they did not appear to alter time to admission to residential care.

The recognition of different behavioral phenotypes for bvFTD is consistent with previous studies. Consensus is lacking in regard to the number of phenotypes; 2 (apathetic and disinhibited), 3 (included a mixed group), or, in the case of our study, 4 phenotypes have been reported. Similarly, 4 neuroanatomic bvFTD phenotypes have been identified in 2 separate studies; however, neither of these investigated specific apathetic or disinhibited behavioral differences. These variations are likely explained by methodologic differences such as our data-driven approach and use of different clinical assessment tools. Other reasons for the divergence in findings may include a different starting point in the analyses. Previous studies set out from neuroanatomic patterns, and some of the earlier studies predate the current bvFTD diagnostic criteria and the delineation of the C9ORF72 gene.

The 4 behavioral phenotypes identified in our study did not differ in terms of disease duration, supporting the existence of distinct behavioral phenotypes beyond what could be explained by behavioral fluctuation across disease progression. Later in progression, however, the distinction becomes less clear, with increasing levels of apathy overall and any expressions of severe disinhibition likely to decline. The outlier from the PSD group provides an example of this: that patient had the longest disease duration and highest levels of apathy in the group and was much more functionally impaired.

Apathy appears to be more common than disinhibition in bvFTD. Indeed, a greater proportion of patients with severe apathy were detected in our study, reflecting previous findings. When present, disinhibition is generally accompanied by apathy. In fact, the presence of severe disinhibition without apathy was rare in our study. This profile may represent a rare phenotype, which warrants further research because it may have implications for clinical management and prognosis. Disinhibition alone did not seem to negatively influence functional impairment in bvFTD, which contrasts with previous findings. This inconsistency is likely to be explained by the very core definition of functional decline and selection of clinical assessments.

Confirmation that severe apathy results in greater functional impairment has substantial clinical relevance for 2 reasons: apathy is pervasive in bvFTD and should be a primary focus of intervention, and disinhibited behaviors (e.g., shoplifting or behaving in a socially inappropriate manner) tend to dominate the clinical picture to such an extent that apathy is often overlooked. Therefore, pharmacologic and nonpharmacologic interventions to improve ADL function (or to delay disease progression because these 2 domains overlap) should emphasize addressing task initiation and planning deficits, which are commonly associated with apathy. Introducing activities that are intrinsically motivating and rewarding, broken down to manageable elements and incorporated into a patient’s routine, has been shown to be of some benefit. Given the differences in functional impairment across these phenotypes, future studies should investigate the implications for caregiver burden. There is potential that caregivers may benefit from targeted approaches according to phenotype.

The 4 bvFTD phenotypes identified in our study had clearly delineated behavioral and functional differences. In contrast, no differences in general cognition were found, similar to a recent study. Overall, other studies demonstrated that patients with a predominantly apathetic phenotype perform more poorly on cognitive assessments than do those with a disinhibited phenotype. Variation in cognitive differences between studies may be down to the choice of general cognitive assessment tools (e.g., Mini-Mental State Examination vs ACE-R) or divergent methods of phenotypic separation (e.g., manual classification vs cluster analysis).

Disinhibited behaviors have been associated with higher levels of caregiver distress, and caregiver distress around neuropsychiatric symptoms can lead to earlier residential care admission. While no differences between phenotypes regarding time to residential care admission were identified in our study, information on the average time from diagnosis to residential care admission for the bvFTD cohort as a whole has clinical relevance. Our result indicating admission to residential care 2.8 years after diagnosis aligns with a recent German study that reported a median of 2.0 ± 2.5 years between diagnosis and residential care admission for a bvFTD cohort. Future studies should investigate time from diagnosis to residential care admission in a larger cohort of bvFTD phenotypes.

The 4 behavioral phenotypes identified in the cluster analysis were further supported by the imaging analyses. Compared to healthy controls, patients with the marked apathy phenotypes (PSA and SA + D) had more extensive brain atrophy than those characterized by mild apathy or marked disinhibition (MA...
with existing literature. This neuroanatomic delineation aligned well with the clinical group differences, which highlights that phenotypes with marked apathy are more functionally impaired than phenotypes with marked disinhibition. This, in turn, is strongly supported by our recent study that demonstrated the critical role of apathy in longitudinal functional decline in FTD.

This study contains some limitations that need to be considered in the interpretation of these results. Foremost is the small size of the PSD subgroup, which has implications for the different group comparisons and survival and imaging analyses. Because of the small sample size of the PSD subgroup and related limited statistical power, imaging analyses were carried out uncorrected at a threshold of \( p < 0.001 \) in comparisons with the control group and at \( p < 0.01 \) in pairwise comparisons between bvFTD phenotypes. This approach, however, yielded more conservative maps of cortical differences than the standard false discovery rate correction for multiple comparisons in patients vs controls. Furthermore, these analyses were constrained with a conservative cluster extent threshold of \( k \geq 50 \text{ mm}^2 \) to balance the risk of type I and type II errors. Finally, we did not have pathologic confirmation of our bvFTD cases. Future studies should address the differences in pathology between these behavioral phenotypes and investigate a larger cohort of the PSD phenotype. In addition, the potential link between bvFTD phenotypes and time to residential care admission needs to be further investigated.

The bvFTD phenotypic categories identified in this study could support clinical decisions on prioritization of interventions to reduce disability and to provide information on prognosis for families. Further potential for in vivo distinction of patients with differing pathologic processes could, in turn, help prevent the dilution of positive effects in therapies that suffering pathologic processes could, in turn, help prevent the dilution of positive effects in therapies. Finally, we did not have pathologic confirmation of our bvFTD cases. Future studies should address the differences in pathology between these behavioral phenotypes and investigate a larger cohort of the PSD phenotype. In addition, the potential link between bvFTD phenotypes and time to residential care admission needs to be further investigated.

The bvFTD phenotypic categories identified in this study could support clinical decisions on prioritization of interventions to reduce disability and to provide information on prognosis for families. Further potential for in vivo distinction of patients with differing pathologic processes could, in turn, help prevent the dilution of positive effects in therapies that occurs when patients are grouped under the single umbrella of one overarching clinical syndrome.

**AUTHOR CONTRIBUTIONS**

C.M. O’Conner: study design and conceptualization, data acquisition, data analysis and interpretation, statistical analysis, drafting and revision of the manuscript. R. Landin-Romero: imaging analysis and interpretation, revision of the manuscript. L. Clemson: study design and conceptualization, revision of the manuscript. C. Kazzik and N. Daveson: data acquisition, revision of the manuscript. J.R. Hodges: study supervision, revision of the manuscript. S. Hsieh: critical revision of the manuscript for intellectual content. O. Piguet: study supervision, study design and conceptualization, revision of the manuscript. E. Mioshi: study design and conceptualization, data acquisition, data analysis and interpretation, study supervision, revision of the manuscript.

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**DISCLOSURE**

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