Which neuropsychiatric and behavioural features distinguish frontal and temporal variants of frontotemporal dementia from Alzheimer’s disease?

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Abstract

Objectives—To investigate the prevalence of changes in mood, personality, and behaviour in frontotemporal dementia (FTD) and Alzheimer’s disease (AD) and hence, which features reliably distinguish between them. To establish whether the frontal and temporal variants of FTD are characterised by different behavioural changes.

Methods—A questionnaire was designed to assess a wide range of neuropsychiatric changes; it incorporated features reported in previous studies of FTD and components of the neuropsychiatric inventory. This was completed by 37 carers of patients with Alzheimer’s disease (AD) and 33 patients with frontotemporal dementia (FTD), comprising 20 with temporal variant FTD (tv FTD) or semantic dementia and 13 with frontal variant FTD (fv FTD). An exploratory principal components factor analysis and discriminant function analysis was applied.

Results—Factor analysis showed four robust and meaningful symptom clusters: factor 1—stereotypic and eating behaviour; factor 2—executive dysfunction and self care; factor 3—mood changes; factor 4—loss of social awareness. Only stereotypic and altered eating behaviour and loss of social awareness reliably differentiated AD from FTD with no effect of disease severity. By contrast, executive dysfunction, poor self care, and restlessness showed a significant effect of disease severity only, with the more impaired patients scoring more highly. Changes in mood were found to be equally prevalent in the three patient groups. Analysis of individual symptoms showed increased rates of mental rigidity and depression in the patients with semantic dementia compared with those with fv FTD. Conversely, the latter group showed greater disinhibition. Discriminant function analysis correctly classified 71.4% overall and 86.5% of the patients with AD.

Conclusions—This questionnaire disclosed striking differences between patients with FTD and AD, but only stereotypic behaviour, changes in eating preference, disinhibition, and features of poor social awareness reliably separated the groups. The patients with fv FTD and semantic dementia were behaviourally very similar, reflecting the involvement of a common network, the ventral frontal lobe, temporal pole, and amygdala. Dys-executive symptoms and poor self care were found to be affected by the severity of the disease, reflecting perhaps spread to dorsolateral prefrontal areas relatively late in the course of both FTD and AD. This questionnaire may be of value in the diagnosis and the monitoring of therapies.

Keywords: frontotemporal dementia; Alzheimer’s disease; semantic dementia; neuropsychiatry

Frontotemporal dementia (FTD) is the term currently used in preference to Pick’s disease to describe a range of non-Alzheimer dementias producing focal lobar atrophy involving the frontal lobe and/or the temporal lobes. Frontotemporal dementia is the second commonest form of primary dementia in the presenium and has an enormous impact on carers as it produces marked changes in personality, behaviour, and communication abilities. Yet, compared to the vast literature on Alzheimer’s disease (AD), there have been relatively few systematic studies of FTD and even fewer have compared substantial groups of patients with FTD and AD.

Although often considered a unitary syndrome, numerous studies have characterised the two major presentations of FTD which reflect the predominant locus of pathology: the progressive change in personality coupled with executive dysfunction associated with the frontal variant of FTD (fv FTD) and the progressive fluent aphasia with breakdown in semantic knowledge (semantic dementia) found in patients with the temporal lobe variant of FTD. Cases with both variants were clearly described by Arnold Pick over 100 years ago, but their delineation and characterisation has occurred only in the past decade or so. In the 1980s workers in Lund and Manchester simultaneously recognised that a significant proportion of patients with dementia presented with a frontal syndrome and coined the term dementia of the frontal type. This was preferred to Pick’s disease as only a minority had specific intraneuronal inclusions (Pick bodies). Other terms used to describe identical cases include frontal lobe degeneration, frontal lobe dementia, and most recently frontal variant FTD.
Neuropsychiatric features of frontotemporal dementia and Alzheimer's disease

Mesulam used the term primary progressive aphasia to refer to patients with a predominant language impairment resulting from focal atrophy. More than 100 patients have now been described with progressive aphasia and although the language disorder is heterogeneous, two fairly distinct syndromes have emerged: progressive non-fluent and progressive fluent aphasia. The latter is also known as semantic dementia as the primary deficit is a profound loss of conceptual knowledge underlying their anoma and progressive loss of verbal and non-verbal comprehension. Patients with semantic dementia have striking focal temporal lobe atrophy and the range of pathological findings are identical to that found in patients with fv FTD. The literature on semantic dementia has focused predominantly on the cognitive and linguistic features, but a recent study suggested that changes in behaviour and personality are also common.

The realisation that patients with identical pathology may present with either frontal or temporal lobe dysfunction, and that both features tend to emerge with time, were factors that led the Lund/Manchester groups to propose the use of the term FTD. After the discovery of tau gene mutations in some familial cases, the use of the term FTD has increased. It is, however, a blanket term which has led to it being treated as a unitary phenomenon rather than recognising the heterogeneity of its presentation. Researchers have tended to amalgamate patients, a process that is likely to blur important distinctions. We think that the cognitive deficits are fundamentally different in patients with frontal pathology and those with predominantly temporal pathology and consider here the issue of whether the behavioural profiles are distinguishable.

A wide range of behavioural abnormalities have been reported in FTD including loss of insight, disinhibition, impulsivity, apathy, reduced empathy for others, poor self care, mood changes, stereotypic behaviour, mental rigidity, and changes in eating patterns. Although there are many good case descriptions of patients with these behavioural changes, the prevalence and, in particular, their specificity to FTD has only just begun to be systematically explored. For instance, Levy et al used a standardised carer interview, the neuropsychiatric inventory (NPI), in an attempt to establish whether the frontal and temporal variants of FTD are characterised by different behavioural changes. Although it is clear that semantic dementia results in profound changes in language and knowledge which do not occur in those with fv FTD, it is unclear whether the two groups differ in the range or frequency of the neuropsychiatric symptoms.

Methods

Patients

Patients were identified through the Memory and Cognitive Disorders Clinic at Addenbrooke's Hospital, Cambridge, England, where they were seen by a senior neurologist (JRH), a consultant psychiatrist, and a clinical neuropsychologist. In addition to a clinical assessment, all patients were given standard psychiatric rating scales, to exclude major functional psychiatric disorders such as depression and schizophrenia. The standard neuropsychological battery applied to all cases included tests of episodic memory (the Warrington recognition memory test, story recall from the WMS-R, and recall of the Rey complex figure); semantic memory (category fluency and naming of 64 items chosen from the Snodgrass and Vanderwart corpus, and 'The pyramids and palm trees test'); visuospatial and perceptual assessments (the visual object and space perception battery); executive function (the modified Wisconsin card sorting test and a letter fluency task); general intellectual abilities (components of the Wais-R); and premorbid IQ (the national adult reading scale). All patients underwent CT or MRI and HMPAO-SPECT, together with the usual battery of screening blood tests to exclude treatable causes of dementia.

All patients have participated in a longitudinal Medical Research Council (UK) funded project investigating aspects of language and memory, and have shown progression of their disease with profiles typical of those reported in the literature.

Three groups of patients were involved in the study: frontal lobe variant FTD (n=13), temporal lobe variant FTD or semantic dementia (n=20), and probable AD (n=37). All patients in the FTD groups fulfilled the recent consensus criteria for frontotemporal lobar degeneration, which recognises the subtypes of frontotemporal dementia, semantic dementia, and progressive non-fluent aphasia. We have chosen to use FTD as the broad superordinate term with subsequent subclassification of patients into fv FTD and semantic dementia.
dementia according to the pattern of neuro-psychological, behavioural, and neuroimaging abnormalities described below:

All patients with fv FTD presented with an informant based history of progressive change in personality and behaviour, with at least five of the following features: loss of insight, disinhibition, apathy, restlessness, emotional lability, distractibility, reduced empathy, impulsivity, social withdrawal, poor self care, and features of Kluver-Bucy syndrome. In addition to these behavioural changes, most of the patients also showed some impairment in executive functioning, as assessed by a verbal fluency task and the modified Wisconsin card sorting test, and difficulties in planning tasks such as the Tower of London test, and subtests from the behavioural assessment of the dysexecutive syndrome. Patients with significant impairment on tests of semantic memory were excluded from this group. Participants with a history of significant head trauma, alcoholism, movement disorder, or any other condition known to impair frontal lobe function were also excluded in line with our locally developed diagnostic criteria for fv FTD.

All patients showed either frontal atrophy on MRI or frontal lobe hypoperfusion on HMPAO-SPECT.

Patients with semantic dementia presented with progressive loss of vocabulary affecting expressive and receptive language in the context of fluent speech production. All patients fulfilled the criteria for semantic dementia previously reported: anomia, impairment in single word comprehension, and impoverished semantic knowledge with relative preservation of phonology, syntax, visuospatial abilities, and day to day (episodic) memory.

In all cases, structural brain imaging by MRI showed focal atrophy involving the polar and inferolateral regions of the temporal lobe. In some cases, the atrophy was clearly bilateral, although in others it was markedly asymmetric, if not unilateral.

The diagnosis of probable AD was made according to the criteria developed by the National Institute of Neurological and Communicative Disorders and Stroke (NINCDS) and the Alzheimer’s Disease and Related Disorders Association (ADRDA), which consist of inclusion and exclusion criteria. All patients presented to the Cambridge Memory Disorders Clinic with a progressive cognitive decline, that was not accounted for by other psychiatric, psychological, behavioural, and neuroimaging abnormalities described below.

Table 1 Demographic variables of the three patient groups

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Age range (SD)</th>
<th>Mean age (SD)</th>
<th>Sex M/F</th>
<th>MMSE range</th>
<th>MMSE mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>Semantic dementia (fv FTD)</td>
<td>20</td>
<td>49–74 (6.25)</td>
<td>63 (6.25)</td>
<td>7/13</td>
<td>3–28</td>
<td>16.4</td>
</tr>
<tr>
<td>fv FTD</td>
<td>13</td>
<td>54–70 (6.03)</td>
<td>60.2 (6.03)</td>
<td>11/2</td>
<td>11–30</td>
<td>24.3</td>
</tr>
<tr>
<td>AD</td>
<td>37</td>
<td>51–82 (6.95)</td>
<td>71.2 (6.95)</td>
<td>19/18</td>
<td>8–30</td>
<td>19.2</td>
</tr>
</tbody>
</table>

Moderate (MMSE<18, n=24). These groups correspond to 0.5, 1, and 2 on the clinical dementia rating scale (CDR). All except three of the patients were living at home, those who were institutionalised had a spouse or relative who still maintained close contact and was therefore able to complete the questionnaire.

The patients with Alzheimer’s disease were significantly older than both FTD groups: (F_{2,55}=14.5, p<0.001, AD>semantic dementia>fv FTD). There were also significant differences for MMSE between patients with semantic dementia and those with fv FTD: (F_{1,47}=3.67, p<0.05, fv FTD>semantic dementia). This is not surprising in view of the known insensitivity of the MMSE to early frontal dysfunction.

Assessment of neuropsychiatric features

A questionnaire was designed following a review of the literature to determine the neuropsychiatric symptoms that are commonly reported in FTD and/or AD. It incorporated questions related to symptoms reported in our previous studies of frontal dementia, components from the neuropsychiatric inventory, and specific questions relating to stereotypic behaviour.

After initial piloting, the final version of the questionnaire consisted of 39 questions investigating the following broad domains: depression, elation, irritability, anxiety, aggression, distractibility, executive functioning, risk taking, empathy, apathy, ritualistic/stereotypic behaviour, aberrant motor behaviour, disinhibition, social withdrawal, hallucinations, delusions, changes in food preference, personal care, and sleep patterns (the full questionnaire is available from the authors).

Carers were asked to rate the frequency of a behaviour on a scale of 0–3. Where a score of zero referred to no change; a score of 1 to occasional occurrence (less than once a week); 2 to a frequent problem (at least once a week); and 3 to a severe behavioural problem. The questionnaire was scored by totalling the carer ratings for all the questions relating to each neuropsychiatric feature; this gave an idea of the prevalence of each of the symptoms.

Test-retest reliability was established by asking 14 of the carers to complete a second questionnaire, 1 week after the first, without reference to it. We found an overall high degree of reliability (r=0.86, p<0.01) which applied to each of the three patient groups: AD (r=0.89, n=7, p<0.01), semantic dementia (r=0.70, n=4, p<0.01), fv FTD (r=0.94, n=3, p<0.01).

Results

The data from the 39 questions were reduced using a principal component factor analysis with varimax factor rotation. This technique is based on the assumption that underlying constructs explain the relation between observed variables.

From an initial scree plot, four principal factors were extracted that best describe the pattern of data collected from these patients (table
The symptoms were considered to be part of the factor if their loading was greater than 0.5. By far the strongest factor (V=10.31), accounting for 26.4% of the variance, which we refer to as stereotypic and eating behaviour, consisted of questions related to ritualised behaviour, clockwatching, stereotyped use of catchphrases, preoccupation with counting and numbers, appetite increase, and changes in food preference towards sweet food.

The remaining three factors were weaker. Factor 2 (V=2.97) accounting for 7.6% of the variance, consisted of questions related to poor judgement, lack of planning, aberrant motor behaviour, distractibility and poor self care which we refer to as executive dysfunction and self care.

Factor 3 (V=2.49) accounting for 6.4% of the variance, lends itself to the term mood changes, revealing heavy loading of questions related to rapid mood changes, depression, and aggression.

Factor 4 (V=2.18) accounting for 5.6% of the variance, we have called loss of social awareness, consisted of questions about disinhibited behaviour, conversational disinhibition, reduction in conversation, and social withdrawal.

There were a few symptoms which seemed to contribute to more than one factor. Lack of empathy, for instance, was just above the cut off at 0.51 in factor 1 and just below, at 0.496, in factor 4. In this case we think that it was more logically in factor 4 along with questions on social withdrawal.

Mental rigidity was just below the cut off for inclusion in factor 1, at 0.49, and just above the cut off in factor 3, at 0.53, but in view of its association with stereotypic and ritualistic behaviour we have included it in factor 1.

Apathy loaded almost equally into factors 2 and 4. Because mood change and cognitive factors are likely to contribute to apathy we have considered this symptom separately.

**INTERGROUP COMPARISONS**

Having established the four principal factors, we then investigated differences between groups on these four measures, using two way analysis of variance (ANOVA) according to diagnosis (AD, semantic dementia, fv FTD) and severity of disease (minimal, mild, and moderate). Specific pairwise comparisons were explored using Tukey’s HSD post hoc test.

Patients who typically showed these behaviours had high, positive scores on the factors, whereas those patients in whom these behaviours were typically absent had low, negative scores.

**Factor 1—Stereotypic and eating behaviour**

Analysis showed a highly significant main effect of diagnosis (F(2,67)=12.6, p<0.001). Post hoc analysis showed significant differences between AD (mean score=−0.59) and both the other groups, but not between fv FTD (mean=0.40) and semantic dementia (mean score=0.83).

Severity of disease was not significant (F(2,67)<1), and there was no interaction between diagnosis and severity (F(4,131)<1).

**Factor 2—Executive dysfunction and self care**

By contrast with factor 1, ANOVA disclosed a significant main effect of severity (F(2,67)=4.48, p<0.05), with significant differences between minimal (mean score=−0.31) and moderate (mean score=0.59), and between mild (mean score=−0.38) and moderate. However, in this case, there was no significant effect of diagnosis (F(2,67)=1.37, p=0.26) and no interaction (F(4,131)<1).

**Factor 3—Mood changes**

For this factor there was no significant effect of diagnosis (F(2,67)<1) or severity (F(2,67)=1.56, p=0.22) and no interaction (F(4,131)<1).

**Factor 4—Loss of social awareness**

As with factor 1, there was a significant effect of diagnosis (F(2,67)=4.50, p<0.05), but in this instance post hoc analysis showed significant differences between fv FTD (mean score=0.60) and AD (mean score=−0.25) only. There was no significant effect of severity (F(2,67)<1) and no interaction (F(4,131)<1).

Although the ANOVAs based upon the factor scores establish the pattern of significant differences between groups, these analyses do not consider the frequency of individual symptoms within each group or which symptoms within the cluster are significantly more common in which group. To explore this important clinical issue, we examined the frequency of key symptoms, using the clusters determined above in the three patient groups. It will be recalled that carers used a four point scale. We decided to include scores of 2 or greater as a positive score. Based on this dichotomy, the frequency of the most important symptoms is shown in the figure.

The symptom which most clearly separated the groups was the use of a catchphrase which occurred in 80% of patients with semantic dementia, 77% of patients with fv FTD, and only 13% of patients with AD. This difference was highly significant (χ² =27.5, p<0.001). Mental rigidity was also extremely common, being present in 80% of the semantic dementia...
patients, 38% of fv FTD, and 27% of AD ($\chi^2=13.5, p<0.01$). Stereotypic/ritualistic behaviour (such as sticking to a very fixed routine or carrying out rituals while doing certain activities) and a preoccupation with counting/numbers were less common, being present in 35% of patients with semantic dementia, 23% with fv FTD, and only 13% with AD ($\chi^2=8.40, p<0.05$).

The symptoms which constituted factor 2; poor planning/judgement, distractibility, aberrant motor behaviour, and poor self care were present in a surprisingly high proportion of all patient groups (figure B). For instance, distractibility was present in 90% of semantic dementia, 69% of fv FTD, and 51% of AD; this difference is significant ($\chi^2=8.69, p<0.05$). Decrease in self care also showed a significant difference, occurring in 69% of fv FTD, 65% of semantic dementia, and only 35% of AD ($\chi^2=6.99, p<0.05$). The only other significant difference in this factor was poor judgement, which was reported in 77% of patients with fv FTD, 65% of semantic dementia, and 40% of AD ($\chi^2=6.42, p<0.05$).

Figure C shows that mood related symptoms, including irritability, were less commonly present and were more or less equally prevalent in all groups, with scores being slightly raised in semantic dementia and fv FTD. The exception

was depression, which was present in 45% of patients with semantic dementia, 24% of patients with AD, and only 7% of the patients with fv FTD ($\chi^2=0.11$).

The prevalence of the symptoms in the final cluster, loss of social awareness was also high across all the groups, but several were present to a higher degree in the FTD subgroups (figure D). Reduction in conversation was present in 95% of patients with semantic dementia, 92% of fv FTD, and 51% of AD; this difference was significant ($\chi^2=15.6$, p<0.01). Social withdrawal occurred in 69% of patients with fv FTD, 60% of those with semantic dementia, and only 27% of those with AD ($\chi^2=10.6$, p<0.05). Similar percentages showed a reduced interest in the family but this difference did not quite reach significance. Disinhibition was fairly prevalent in the fv FTD group, 46% compared with just 5% of the semantic dementia group and the AD group. Significant differences were found in both conversational disinhibition ($\chi^2=17.7$, p<0.001) and disinhibited behaviour ($\chi^2=7.65$, p<0.05). These two questions have been combined on the graph to give an overall measure of disinhibition.

Delusions were reported in 15% of patients with fv FTD, 11% with AD, and only 5% with semantic dementia (figure E). Hallucinations were even less commonly reported and occurred in 15% of patients with AD, 8% of patients with fv FTD, and none of the patients with semantic dementia. None of these differences were significant.

Apathy was reported in 73% of patients with fv FTD, 65% of semantic dementia, and only 54% of AD. This difference was not significant.

**DISCRIMINANT ANALYSIS**

A stepwise discriminant function analysis was performed to determine which symptom clusters differentiated the patient groups with the highest diagnostic accuracy. Factor 1—stereotypic and eating behaviour—was shown to discriminate between the patient groups most reliably, followed by factor 4—loss of social awareness. It is not surprising, following the results of the ANOVAs, that factors 2 and 3—executive dysfunction and self care and mood changes—did not contribute to the discriminant analysis.

Overall, 71.4% of patients were correctly classified, 55% of patients with semantic dementia, 86.5% of patients with AD, and 53.8% of patients with fv FTD. The patients with fv FTD tended to be misclassified as semantic dementia (38.5%), with only a small proportion being misclassified as AD (7.7%). Of the patients with semantic dementia, 20% were misclassified as AD and 25% as fv FTD. Only a small proportion of patients with AD were not correctly classified and they were equally misclassified as semantic dementia (5.4%) and fv FTD (8.1%).

**Discussion**

As predicted, our behavioural questionnaire disclosed striking differences between patients with FTD and AD, although the number of domains in which such differences existed was perhaps less than expected from published literature. Of the four symptom clusters disclosed by factor analysis only two, stereotypic and eating behaviour (factor 1) and loss of social awareness (factor 4), were significantly more common in the FTD group. By far the strongest factor was the former, accounting for 26% of the overall variance. Of the symptoms within this cluster, the use of a catchphrase, mental rigidity, and alteration in appetite were by far the most discriminating. Within factor 4, reduction in conversation, social withdrawal, loss of interest in family, and disinhibition were the principal symptoms which separated FTD and AD. It is interesting to note that neither mood nor dysexecutive symptoms distinguished the groups. In previous studies many symptoms have been listed as characteristic of FTD. It is clear, however, from this study that only certain symptoms clearly distinguish FTD from AD.

Rather surprisingly, we found no significant differences between patients with semantic dementia and fv FTD using the factor analysis. Examination of the proportion of patients scoring highly on the individual symptoms showed, however, definite but subtle differences. The patients with semantic dementia scored much more highly on mental rigidity and depression than the patients with fv FTD. Conversely, patients with fv FTD were more disinhibited.

We were particularly interested in the range and frequency of stereotypical behaviour in FTD and included a wider range of questions than previous workers. We found routine, mental rigidity, clockwatching, use of a catchphrase, ritualised behaviour, and a preoccupation with counting and numbers to be increased in both of the FTD groups compared with AD. Only factor 2—executive dysfunction and self care—showed an effect of severity of disease, with the more impaired patients scoring more highly on these questions.

Overall, our results support the findings of Miller et al in their study evaluating the Lund/Manchester criteria. They found 12 out of the 17 symptoms that they investigated to be significantly higher in patients with FTD than in patients with AD. Eleven of these symptoms were included in our questionnaire and we found the same significant differences except for distractibility and impulsivity, which were equally prevalent in FTD and AD. Miller et al reported that loss of personal awareness, eating, perseverative behaviour, and a reduction in speech most clearly differentiated FTD from AD, three out of four of these symptoms were included in our questionnaire (not loss of personal awareness) and were found to be significantly increased in the FTD group.

Like Levy et al, we found patients with FTD to show increased levels of disinhibition and lower levels of depression. Apathy, euphoria, and aberrant motor behaviour were, however, equally prevalent in all of our patient groups. Lebert et al, using the frontotemporal behavioural scale, found increased scores in FTD on all four of their symptom clusters; self
monitoring, self neglect, self centred behaviour, and mood changes. Our results support these findings with the exception of mood symptoms, which were found to be equally common in both FTD and AD. Depression was only reported in one of our patients with fv FTD.

Of the symptoms in our study that overlap with those in the frontal behavioural inventory, only disinhibition was found to be raised in FTD compared with AD. In a pilot study using this instrument, Kertesz et al found higher levels of disinhibition, euphoria, aggression, and perseveration in FTD.

It is clear, therefore, that patients with fv FTD and semantic dementia are behaviourally very similar and that the main distinction is that the second results in marked semantic deficits. The high prevalence of behavioural changes in semantic dementia may have been overlooked in the past because of the profound linguistic deficits which interfere with patient evaluation. It is also possible that the behavioural symptoms occur at a later stage in semantic dementia.

Turning to the neuroanatomical implications of our findings, it is likely that the behavioural syndrome reflects the involvement of a common network in both variants, namely, the ventral (orbitobasal) frontal lobe, temporal pole, and amygdala. The two clusters of symptoms which distinguished FTD from AD, stereotypic behaviour, change in food preference, and loss of social awareness have all been linked to ventral frontal pathology. It seems likely that this region is affected from an early stage in all patients with both variants, either by direct pathological involvement or indirectly via damage to the temporal pole and amygdala, which are heavily interconnected with the ventromedial frontal lobe. The ventral frontal cortex has long been associated with regulating inhibitory control and damage leads to a failure to inhibit inappropriate responses. Based on these well established findings, Piaisted and Sahakian proposed that deficits in social cognition in patients with frontal lobe damage are due to an inability to inhibit inappropriate behaviours. They argued that this prevents the selection of more appropriate action plans dictated by long term goals. An alternative explanation comes from Damasio et al who proposed that damage to the ventromedial frontal cortices leads to an inability to activate somatic states linked to punishment and reward previously associated with specific social situations. Normally, these are activated in connection with anticipated outcomes of response options. A third theory for the breakdown in social behaviour is the “theory of mind” view, which suggests that these patients have lost the ability to make inferences about the mental states of others.

This view is supported by the finding that these patients do show a loss of empathy for others, and observations that conversation becomes increasingly self centred.

Edwards Lee et al reported many of the same behavioural deficits in temporal lobe variant FTD and suggested that either the temporal lobes themselves are important mediators of social behaviour, or the ventromedial frontal cortex and anterior temporal structures work reciprocally. There is strong evidence from both human and monkey studies to support the second view. Medial temporal lobe lesions affecting the limbic system including the amygdala have long been associated with behavioural deficits. Ablation studies in monkeys have shown that damage to the ventromedial frontal cortex leads to disturbances of emotional and social behaviour.

In humans, the ventromedial frontal cortex has also been implicated in decision making, risk assessment, and in pathophysiology of obsessive compulsive disorders. In keeping with these findings Mummery et al suggest that behavioural disturbances in FTD arise from disruption to a network of limbic regions, including the ventromedial frontal cortex, amygdala, and connections to anterior temporal lobe structures.

The differences in behavioural profiles between patients with the two variants of FTD are also of potential interest. Patients with fv FTD showed higher levels of disinhibition and very low reports of depression, whereas patients with semantic dementia showed greater attentional and memory deficits. The fv group experience a gradual loss of expressive and receptive vocabulary; clinging to a fixed routine and becoming extremely rigid in their thinking may, therefore, be an attempt to maintain some control and understanding over a world that is becoming increasingly incomprehensible.

By contrast with the clear separability of the groups by symptoms related to putative ventromedial frontal pathology, it is interesting that we found an effect of severity only for factor 2 (dysexecutive function and self care). Dysexecutive ability reflects dorsolateral prefrontal functioning and this region is involved relatively late in the course of AD, when the disease has spread from the hippocampal and posterior association cortex. We hypothesise that in FTD, pathology also spreads upwards to involve this region late in the course of the disease. Decrease in levels of self care could reflect problems with planning and organisation, or might be considered to be a consequence of general apathy.

Changes in mood, except for depression, were equally common in all three patient groups and seemed, therefore, to be of limited diagnostic value. Depressed mood may occur as a psychological response to a diagnosis of a neurodegenerative disease, or it may be caused by changes in the neurochemical systems. The lack of depression found in patients with fv FTD in this study accords with previous findings and may reflect the almost universal lack of insight found in these patients.

Apathy was commonly reported in all patient groups, with scores being higher in semantic dementia and fv FTD, although the prevalence was rather less than previously reported in both AD and FTD. It is also surprising that delusions and hallucinations were present in such a small minority of patients from all three groups. Many studies have looked at the preva-
lence of these symptoms in AD and the results have varied widely. Delusions have been found to affect 10% to 75%, and hallucinations 3% to 40%. Our results are more in keeping with those of Cummings who found delusions to affect 22% of patients and hallucinations 10% of patients with AD.

Our study has, of course, some potential shortcomings. Firstly, many of the patients included were quite late in the course of their disease. We attempted to compensate for this by the segregation of cases by disease severity, although the MMSE is not a very satisfactory instrument for this purpose in FTD. In future work, it would obviously be desirable to include more cases newly presenting to the clinic. Secondly, it might be argued that a study of this type entails a degree of circularity in that the patients with fv FTD were selected, partly, on the basis of prominent behavioural symptoms (in addition to radiological and neuropsychological features). It should be emphasised, however, that the features found to discriminate FTD from AD were a specific subset of a wider range used to select cases. Moreover, the patients with semantic dementia were selected on the basis of cognitive and neurological features, yet were found to have a high prevalence of neuropsychiatric symptoms. Finally, any study of this type must be regarded as preliminary, our findings require confirmation in a new prospectively assessed cohort.

Overall, our questionnaire correctly classified 71.4% of the patients involved in this study and only 13% of patients with AD were incorrectly classified. It is hoped that it will prove to be a useful tool for discriminating between the dementias both in a clinical and research setting. This has become increasingly important as therapies specifically targeted at AD have emerged and are sought for FTD. Changes in personality and behaviour are often the earliest symptoms of dementia of the frontal type and are usually the most distressing aspect of the disease for the carers. At presentation, these patients may perform flawlessly on conventional cognitive assessments, it is, therefore, imperative to find ways of accurately quantifying neuropsychiatric features if we are to be able to diagnose patients early in the course of the disease.

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We are indebted to the continuing support of the patients and carers included in this study. We are also very grateful to Peter Watson for his statistical advice. This work was supported as part of a Medical Research Council (UK) programme grant to RH. MALR is supported by a National Institutes of Health V(USA) grant.


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Sasha Bozeat, Carol A Gregory, Matthew A Lambon Ralph and John R Hodges

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