Structural neuroanatomy of face processing in frontotemporal lobar degeneration

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ABSTRACT

Impairments of face processing occur frequently in frontotemporal lobar degeneration (FTLD) but the neuroanatomical basis for these deficits has seldom been studied systematically. Here a prospective voxel based morphometry study is described addressing the neuroanatomy of two key dimensions of face processing—face identification and facial emotion recognition—in a single cohort of 32 patients with FTLD (19 with frontal variant and 13 with temporal variant FTLD). For the FTLD group as a whole, face identification was positively associated with grey matter in the right anterior fusiform gyrus while recognition of angry expressions was positively associated with grey matter in the bilateral insula cortex. FTLD provides a perspective on the neuroanatomy of face processing that is complementary to focal lesion and normal functional imaging work.

INTRODUCTION

Frontotemporal lobar degeneration (FTLD) is a clinically and pathologically diverse group of neurodegenerative diseases associated with focal frontal and temporal lobe atrophy. Behavioural disturbances are common in FTLD, most saliently in the canonical syndromes of behavioural variant frontotemporal dementia (bvFTD) and semantic dementia (SD). Behavioural change may be partly attributable to defective processing of social signals: in particular, impaired processing of information about human faces is an important clinical issue in both bvFTD and SD. However, the nature of this impairment is likely to differ between the syndromes. SD is characterised by panmodal breakdown of knowledge about objects, concepts and persons, leading to impaired processing of facial identity; progressive prosopagnosia has been associated particularly with selective right temporal lobe atrophy. In contrast, bvFTD is characterised by decline in interpersonal and executive skills, with loss of emotional responsivity and impaired recognition of emotional facial expressions. This differential involvement of face processing functions may assist in early clinical diagnosis and differentiation of SD and bvFTD. Anatomically, the frontal and temporal cortical areas affected in FTLD overlap brain regions implicated in face processing from focal lesion studies and functional imaging work in healthy subjects, including the anterior and inferior temporal lobes. However, few studies have addressed systematically the neuroanatomical basis of face processing deficits in FTLD using unbiased techniques such as voxel based morphometry (VBM).

Here we investigated prospectively the structural neuroanatomical substrates of two key dimensions of face processing—identification of familiar faces and recognition of facial emotions—using VBM in a single cohort of patients with FTLD. Based on previous work in patients with focal brain lesions and in healthy subjects, we hypothesised that face identification would correlate with grey matter in inferior temporal regions while processing of facial emotions would correlate with grey matter in temporal, inferior frontal and insula regions, with partially separable anatomical associations for different facial emotions.

METHODS

Thirty-two patients fulfilling consensus clinical criteria for FTLD were recruited. All patients had a general neuropsychological assessment and volumetric brain MRI. The study cohort comprised 19 patients with predominant frontal lobe atrophy (frontal variant (fv) FTLD), each fulfilling clinical criteria for bvFTD; and 13 patients with predominant asymmetric (left or right) temporal lobe atrophy (temporal variant (tv) FTLD), 12 fulfilling clinical criteria for SD (semantic progressive aphasia) and one (with predominantly right-sided temporal lobe atrophy) presenting with progressive prosopagnosia. Patients were compared neuropsychologically with a group of 22 age matched healthy controls recruited via the Dementia Research Centre database. Subject characteristics are summarised in table 1. Informed consent was obtained in each case and the study was approved by the local research ethics committee in accord with the Declaration of Helsinki guidelines.

All patients were assessed on tests of face identification, affective processing and perceptual analysis (table 1). Face identification was assessed using a Famous Faces Test requiring recognition of black and white photographs of 12 contemporary public figures widely familiar to UK residents (regularly updated from Warrington and James). Identities of the individuals in the test (comprising six prominent British politicians, two prominent American politicians, two high profile entertainers and two members of the British Royal Family) are listed in supplementary table 1 (available online only). Evidence of recognition other than naming can be used to score performance on this test, making it suitable for use in individuals with naming impairments (eg, in SD). Facial emotion recognition was assessed using 24 faces from the Ekman and Friesen battery. Perceptual analysis of faces was assessed using the Benton Facial Recognition Test. Details of all neuropsychological tests used are provided in the supplementary material (available online only).
Statistical parametric maps (SPMs) for each behavioural contrast were assessed at the whole brain level and after small volume correction using the previously defined anatomical regions at family wise error (FWE) corrected threshold p<0.05.

RESULTS

Neither the tvFTLD nor the fvFTLD subgroup (analysed separately) showed a VBM association of face or emotion identification performance at the whole brain level or following small volume correction. For the combined FTLD group, SPMs of regional grey matter positively associated with performance on the face and emotion identification tasks are displayed in figure 1 (regional maxima for these SPMs are summarised in supplementary table 2, available online only). Performance on famous face identification was associated with grey matter volume in the right anterior fusiform gyrus (p<0.05 after FWE correction within the prespecified anatomical small volume). Performance on recognition of anger was associated with grey matter volume in the posterior insula cortex bilaterally (p<0.05 after FWE correction within the prespecified anatomical small volume). No other anatomical associations of face identification or individual emotion recognition and no anatomical associations of performance on the face perception task were identified after whole brain or small volume correction at the prescribed threshold. Direct structural contrasts between the tvFTLD and fvFTLD subgroups using VBM did not demonstrate any grey matter voxels overlapping with the anatomical associations of face or emotion identification (even at a lenient statistical threshold of p<0.001 uncorrected), suggesting that the behavioural regressions were not driven simply by anatomical differences between subgroups.

DISCUSSION

Here we have demonstrated structural neuroanatomical substrates for two key dimensions of face processing in a cohort of patients with FTLD. The findings corroborate previous evidence for deficient identification of faces and facial expressions in FTLD\(^7\) and show that these different components of face processing depend on distinct brain areas. Face identification was associated with grey matter volume in the right fusiform gyrus, somewhat anterior to the fusiform face area (the cortical area previously emphasised as a key site for the abstraction of facial identity information based on focal lesion and normal functional imaging evidence\(^4\)). It is unlikely we have demonstrated an anatomical association of face perception per se as the association with face identification performance was present after adjusting for purely perceptual performance. We suggest that this more anterior inferior temporal region linked with face

![Table 1](http://jnnp.bmj.com/)
identification may be involved in semantic memory for faces, consistent both with previous studies of associative prosopagnosia in FTLD\textsuperscript{3} \textsuperscript{13} and with functional imaging studies of familiar face processing in normal individuals.\textsuperscript{7}

In addition, we have shown an anatomical association for recognition of angry facial expressions in the insula cortex. Anger recognition has previously been identified as a key deficit in FTLD.\textsuperscript{6} Information about brain regions critical for anger recognition is limited but the insula has been implicated previously in the processing of anger by the healthy brain.\textsuperscript{8} The lack of a grey matter association for recognition of other facial expressions may at first appear somewhat surprising in light of present and previous behavioural findings in FTLD.\textsuperscript{5,6} However, whereas anger is a basic facial expression posing a potential threat to self, the relatively greater conceptual complexity involved in interpreting other facial emotions, such as happiness, fear or disgust,\textsuperscript{14} might make detection of a consistent group level VBM association more difficult. Moreover, VBM work in Huntington’s disease has demonstrated an association between insula volume and disgust (but not anger) recognition\textsuperscript{15}; this suggests that the neuroanatomical profile of facial emotion processing may be modulated by the underlying neurodegenerative pathology, and further supports functional imaging evidence in normal subjects that the insula is engaged in processing various emotions according to task demands.\textsuperscript{9} It is also noteworthy that the amygdala was not identified as an association of emotion recognition in our study or in a previous VBM study of FTLD,\textsuperscript{5} which may reflect the relatively low behavioural value and limited arousal potential of the face stimuli for these FTLD patients.

Neuroanatomical associations in a diffuse neurodegenerative disease such as FTLD should be interpreted with care, particularly where (as here) small case numbers limit the strength of those associations. However, this study suggests that neuroanatomical findings concerning face processing mechanisms in FTLD complement focal lesion studies and normal functional imaging work. Like focal lesion studies (and in contrast to normal functional imaging studies), the evidence from neurodegenerative disease can identify brain regions that are critical for particular aspects of face processing. The present findings support cognitive models that posit dissociable processing modules for different aspects of face analysis.\textsuperscript{16} In future work it will be important to study face processing mechanisms using combined behavioural and neuroimaging approaches in a range of neurodegenerative diseases, and to assess the specificity of these mechanisms for processing faces versus other stimuli of comparable perceptual and semantic complexity.

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Competing interests None.

Ethics approval This study was conducted with the approval of the the National Hospital for Neurology and Neurosurgery and Institute of Neurology Joint REC.

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