Voice processing in dementia: a neuropsychological and neuroanatomical analysis

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Voice processing in neurodegenerative disease is poorly understood. Here we undertook a systematic investigation of voice processing in a cohort of patients with clinical diagnoses representing two canonical dementia syndromes: temporal variant frontotemporal lobar degeneration (n = 14) and Alzheimer’s disease (n = 22). Patient performance was compared with a healthy matched control group (n = 35). All subjects had a comprehensive neuropsychological assessment including measures of voice perception (vocal size, gender, speaker discrimination) and voice recognition (familiarity, identification, naming and cross-modal matching) and equivalent measures of face and name processing. Neuroanatomical associations of voice processing performance were assessed using voxel-based morphometry. Both disease groups showed deficits on all aspects of voice recognition and impairment was more severe in the temporal variant frontotemporal lobar degeneration group than the Alzheimer’s disease group. Face and name recognition were also impaired in both disease groups and name recognition was significantly more impaired than other modalities in the temporal variant frontotemporal lobar degeneration group. The Alzheimer’s disease group showed additional deficits of vocal gender perception and voice discrimination. The neuroanatomical analysis across both disease groups revealed common grey matter associations of familiarity, identification and cross-modal recognition in all modalities in the right temporal pole and anterior fusiform gyrus; while in the Alzheimer’s disease group, voice discrimination was associated with grey matter in the right inferior parietal lobe. The findings suggest that impairments of voice recognition are significant in both these canonical dementia syndromes but particularly severe in temporal variant frontotemporal lobar degeneration, whereas impairments of voice perception may show relative specificity for Alzheimer’s disease. The right anterior temporal lobe is likely to have a critical role in the recognition of voices and other modalities of person knowledge.

Keywords: voice; face; phonagnosia; frontotemporal dementia; Alzheimer’s disease
Abbreviations: FTD = frontotemporal dementia; FTLD = frontotemporal lobar degeneration
Introduction

Disorders of face processing are well recognized and widely studied, but much less attention has been paid to disorders of voice processing. From a neurobiological perspective this is somewhat surprising, since human voices can be considered ‘auditory faces’ in several important respects (Schweinberger et al., 1997a; Belin et al., 2004). Ethologically, voices are the second major channel via which we identify other people, while perceptually, individual voices (like faces) present the brain with a formidable problem of precise differentiation and identification within a class of complex sensory objects. The relative neglect of voice processing in the clinical literature is likely to reflect a lack of theoretical frameworks for understanding voice analysis, the comparative difficulty of working with vocal stimuli and the scarcity of symptomatic deficits of voice processing (phonagnosia), in turn partly attributable to the primacy of the face as a source of person knowledge in daily life. However, this situation has been transformed recently with the development of cognitive models of voice processing (Belin et al., 2000, 2004; von Kriegstein et al., 2005, 2006; Campanella and Belin, 2007; Goll et al., 2010), the availability of sophisticated stimulus synthesis and delivery systems, functional imaging studies in the healthy brain and the detailed characterization of phonagnosias. Cognitive models of voice processing have been influenced by face processing models (Bruce and Young, 1986; Burton et al., 1990; Ellis et al., 1997). In the voice processing model proposed by Belin et al. (2004), voice identification occurs via serially and hierarchically organized processing stages: perceptual representations or templates derived from modality-specific (voice and face) recognition units feed into amodal ‘person identity nodes’, evoking a sense of familiarity that gates access to biographical information (including personal names) and leads to recognition of the speaker. These putative processing stages map onto an anatomical hierarchy in which perceptual analysis of voices occurs in temporoparietal cortices with associative processing of voices and other modalities of person knowledge in more anterior temporal lobe areas, delineated using functional brain imaging (Imaizumi et al., 1997; Nakamura et al., 2001; Belin et al., 2002; von Kriegstein et al., 2006; Warren et al., 2006; Bishop and Miller, 2009). However, the cognitive and neural architecture of voice processing and its relations to other modalities of person knowledge continue to be defined. Phonagnosia has been described as a developmental disorder (Garrido et al., 2009) and, more commonly, in association with focal damage involving the right or left temporal lobe or the right parietal lobe (Van Lancker and Canter, 1982; Van Lancker and Kreiman, 1987; Van Lancker et al., 1988, 1989; Ellis et al., 1989; Hanley et al., 1989; Neuner and Schweinberger, 2000; Belin et al., 2004; Lang et al., 2009).

There are both clinical and neurobiological grounds for a systematic analysis of voice processing in the degenerative dementias. Clinically, voice processing impairments are likely to be under-recognized in these diseases yet may constitute a significant and disabling symptom (Hailstone et al., 2010), especially in situations where additional cues to speaker identity are reduced or unavailable. Neurobiologically, the common dementias collectively affect distributed areas in the temporal, parietal and frontal lobes that have been implicated in voice processing in functional imaging studies of healthy subjects (Seeley et al., 2009). In particular, the brunt of tissue damage in Alzheimer’s disease and several diseases in the frontotemporal lobar degeneration (FTLD) spectrum initially falls on the temporal lobes, which are likely to contain mechanisms integral for voice analysis and recognition (Belin et al., 2004). The syndrome of progressive prosopagnosia is well recognized in association with right temporal lobe atrophy (Evans et al., 1995; Joubert et al., 2003, 2004; Josephs et al., 2008; Chan et al., 2009) and is dominated by deficits of non-verbal knowledge, in particular knowledge of familiar people (Hanley et al., 1989; Gentileschi et al., 1999, 2001; Snowden et al., 2004; Thompson et al., 2004; Gainotti, 2007, 2008). While voice recognition commonly becomes affected with evolution of the progressive prosopagnosia syndrome (Gentileschi et al., 2001; Gainotti et al., 2003, 2008), selective phonagnosia has seldom been reported. However, progressive associative phonagnosia with relatively-preserved voice discrimination, face and proper name recognition has recently been described in association with frontotemporal atrophy involving the right anterior temporal lobe (Hailstone et al., 2010). Neurodegenerative diseases offer a perspective on voice processing that is complementary both to studies in normal subjects and in patients with focal brain lesions: the breakdown of voice processing in dementia would potentially allow identification of critical nodes in a functional and anatomical cerebral network, and inform cognitive models of voice processing.

In this study, we investigated neuropsychological and neuroanatomical signatures of voice processing in two canonical dementias, FTLD and Alzheimer’s disease. In targeting these disease groups we recognized that, whereas Alzheimer’s disease typically presents with a relatively uniform clinical and anatomical profile, FTLD is clinically and anatomically heterogeneous. In particular, patients with FTLD who have predominant temporal lobe atrophy (i.e. those predicted a priori to develop voice processing deficits) have heterogeneous clinical presentations, including both semantic dementia (progressive semantic aphasia or progressive prosopagnosia) and progressive behavioural decline (behavioural variant frontotemporal dementia (FTD)). For the purposes of the present ‘lesion-led’ study, we selected patients with FTLD based on the presence of predominant temporal lobe atrophy: we term this non-canonical, anatomically defined subgroup ‘temporal lobe variant FTLD’ (Brambati et al., 2009). In line with current cognitive models and previous neuropsychological evidence concerning the organization of voice processing (Van Lancker et al., 1988; Ellis et al., 1997; Schweinberger et al., 1997b; Neuner and Schweinberger, 2000; Belin et al., 2004; Lucchelli and Spinelli, 2008; Garrido et al., 2009; Hanley and Damjanovic, 2009), we designed a series of neuropsychological experiments incorporating subtests to assess early perceptual encoding, discrimination and recognition of voices in the target clinical groups. Processing of voices was assessed in relation to processing of faces and names, in order to assess the modality- and material-specificity of any voice processing deficit. Neuroanatomical correlates of voice processing performance were assessed using voxel-based morphometry (Ashburner and Friston, 2000). We hypothesized distinct
profiles of phonagnosia in temporal lobe variant FTLD and Alzheimer’s disease, with more severe associative impairment in temporal lobe variant FTLD and relatively more prominent apperceptive impairment in Alzheimer’s disease. We further hypothesized, based on anatomical evidence in the healthy brain, that semantic deficits in processing voices (in common with other kinds of person knowledge) would be associated with atrophy of anterior temporal lobe regions, and voice apperceptive deficits would be associated with atrophy of more posterior temporoparietal regions.

Materials and methods

Subject details

Fourteen consecutive patients with temporal lobe variant FTLD, 22 patients with Alzheimer’s disease and 35 healthy older control subjects participated. All patients were recruited from the tertiary Cognitive Disorders Clinic at the National Hospital for Neurology and Neurosurgery. Patients with temporal lobe variant FTLD were selected based on the presence of selective, bilateral anterior temporal lobe atrophy on MRI (atrophy of one or both temporal lobes disproportionate to any accompanying atrophy of other cerebral regions, as assessed visually by an experienced, independent neuroradiologist); the distribution of temporal lobe atrophy was asymmetric in 13/14 cases. Clinically, most (13/14) patients with temporal lobe variant FTLD had a syndrome of semantic dementia according to the consensus criteria of Neary et al. (1998); within this semantic dementia subgroup, 10 patients presented with progressive semantic aphasia and three presented with progressive prosopagnosia. One patient within the FTLD group presented with behavioural variant FTD. All patients with Alzheimer’s disease had a clinical syndrome of typical Alzheimer’s disease led by memory decline and fulfilled modified National Institute of Neurological and Communicative Diseases and Stroke/Alzheimer’s Disease and Related Disorders Association (NINCDS-ADRDA) criteria for probable Alzheimer’s disease (Dubois et al., 2007). Twenty out of 22 patients with a clinical diagnosis of Alzheimer’s disease had brain MRI: 16 patients had disproportionate symmetrical hippocampal atrophy and four had generalized cerebral atrophy. Demographic characteristics of the study groups are summarized in Table 1 and further details are provided in the online Supplementary material. Significance of differences between groups was assessed using z-tests with bootstrap (2000 replicates) standard errors. Subject groups did not differ significantly in age, gender distribution or years of education (all

Table 1 Summary of subject characteristics

|                            | Temporal lobe variant FTLD n = 14 | Alzheimer’s disease n = 22 | Healthy controls n = 35 |
|-----------------------------|-----------------------------------|-----------------------------|-------------------------|
| Demographic characteristics |                                    |                             |                         |
| Males: females              | 8:6                               | 10:12                       | 13:22                   |
| Right: left-handed          | 11:3                              | 19:3                        | 31:4                    |
| Age (years)                 | 64.2 (6.3)                        | 66.5 (7.7)                  | 63.9 (5.7)              |
| Years of education          | 13.9 (4.8)                        | 13.5 (3.6)                  | 15.2 (3.3)              |
| Clinical characteristics    |                                    |                             |                         |
| Clinical syndrome at presentation | Semantic dementia (n = 13)
| Behavioural variant FTD (n = 1) | 5.4 (1.7)                    | 5.7 (2.4)                  | 23.1 (4.2)               |
| Symptom duration (years)    | 3–8                               | 2–11                        | 29.4 (0.6)
b | 28–30                   |
| Mini-Mental State Examination (/30) | 21.1 (7.2)**                      | 21.3 (4.2)** **            | 18d                     |
| Medication                  | 4c                                |                             |                         |
| Cardinal symptoms           |                                    |                             |                         |
| Voice recognition           | n = 9 (64%)                       | n = 11 (50%)                | NA                      |
| Face recognition            | n = 7 (50%)                       | n = 8 (36%)                 | NA                      |
| Voice familiarity           | n = 3 (21%)                       | n = 3 (14%)                 | NA                      |
| Face familiarity            | n = 5 (36%)                       | n = 2 (9%)                  | NA                      |
| Media exposure              |                                    |                             |                         |
| TV watching (hours per week)| 15.1 (9.2)                        | 15.9 (10.0)                 | 14.4 (10.5)             |
| Radio listening (hours per week) | 2.4 (4.2)**‡                     | 11.4 (13.3)                 | 13.8 (12.0)             |
| News exposure (times per week) | 8.1 (4.3) **†                     | 13.0 (8.0)                  | 13.9 (6.9)              |

Assessed using questionnaire (Supplementary material).

a Ten cases with progressive semantic aphasia, three cases with progressive prosopagnosia.

b Twenty-three controls performed a Mini-Mental State Examination.

c Two patients taking a serotonin reuptake inhibitor, one taking anti-Parkinson’s medication, one taking lithium.

d Sixteen patients taking a cholinesterase inhibitor, two taking memantine.

e Assessed using questionnaire (Supplementary material).

NA Not applicable to controls.

*Significantly worse than controls (P < 0.05); **significantly worse than controls (P < 0.001); †significantly different from other patient group (P < 0.05); ‡significantly different from other patient group (P < 0.01).
$P > 0.05$; the temporal lobe variant FTLD and Alzheimer’s disease groups did not differ significantly on two general measures of clinical severity (symptom duration and Mini-Mental State Examination score). The study was approved by the local institutional research ethics committee and all subjects gave informed consent in accord with the principles of the Declaration of Helsinki.

### General neuropsychological assessment

All patients and 19 healthy control subjects had a comprehensive general neuropsychological assessment; the remaining 16 control subjects performed a reduced set of tests. The tests administered are listed in Table 2. Fisher’s exact test was used to assess group differences in gender, for all other variables differences in means were assessed using z-tests with bootstrap (2000 replicates) standard errors.

### Peripheral hearing assessment

Most subjects had no clinical history of hearing loss. To assess any effects of hearing loss on performance in the experimental tasks across the experimental groups, all subjects underwent pure tone audiometry on frequencies between 0.5 and 4 kHz. The audiometry procedure and analysis are described in the online Supplementary material.

### Experimental behavioural investigations: perceptual analysis of voice attributes

Voices are complex auditory objects and voice perception depends on encoding spectro-temporal attributes that carry information about vocal identity such as gender and size (Belin et al., 2002, 2004; Griffiths and Warren, 2004; Warren et al., 2006). Here we created novel subtests to assess encoding of the voice attributes of gender and vocal tract length (an index of vocal size).

### Vocal gender

Vocal gender is determined by lower level perceptual properties including pitch and vocal tract length; here we were interested in subjects’ ability to assign gender to vocal samples based on all available auditory

| Test (max score) | Temporal lobe variant FTLD n = 14 | Alzheimer’s disease n = 22 | Healthy controls n = 35 |
|-----------------|----------------------------------|---------------------------|------------------------|
| WASI Verbal IQ   | Mean (SD) Range                   | Mean (SD) Range           | Mean (SD) Range        |
| WASI Performance IQ | Mean (SD) Range               | Mean (SD) Range           | Mean (SD) Range        |
| Reading IQa      | Mean (SD) Range                   | Mean (SD) Range           | Mean (SD) Range        |
| Semantic tests   |                                  |                           |                        |
| BPVS (/150)      | 73.8 (49.7)**†                   | 141.4 (11.9)*              | 148.1 (1.5) 144–150    |
| Concrete synonyms (/25) | 14.2 (5.3)**†               | 20.9 (2.7)**              | 24.3 (1.3) 19–25       |
| Abstract synonyms (/25) | 15.4 (5.8)**†               | 20.9 (3.5)**              | 24.3 (1.2) 20–25       |
| Landmark name (/15) | 2.6 (3.7)**†                   | 6.1 (4.0)**               | 13.5 (1.3)d 11–15      |
| Landmark identify (/15) | 4.6 (4.7)**†                | 8.0 (4.1)**               | 13.7 (1.2)d 11–15      |
| Other non-semantic skills |                                  |                           |                        |
| GNT (/30)        | 2.2 (6.1)**†                    | 11.6 (7.9)**              | 26.0 (2.4) 19–30       |
| ODT (/20)        | 16.5 (5.0)                      | 15.8 (2.8)**              | 18.5 (1.2) 16–20       |
| Forwards DS (/12) | 7.3 (2.7)                     | 7.1 (2.3)*                | 8.7 (2.0) 4–12         |
| Reverse DS (/12) | 6.1 (3.3)                       | 4.9 (2.7)*                | 7.4 (2.6) 2–12         |
| GDA (/24)        | 8.9 (6.9)**                     | 5.5 (4.5)**               | 15.4 (4.8)c 6–23      |
| Episodic memory  |                                  |                           |                        |
| RMT words (/50)  | 35.4 (7.0)**†                   | 30.1 (7.3)**†             | 47.3 (1.8)a 43–49     |
| RMT faces (/50)  | 28.9 (4.1)**†                   | 35.0 (5.6)**              | 42.2 (4.7)c 35–49     |
| Executive function |                                  |                           |                        |
| Stroop word reading: scaled score | 5.2 (4.0)**†                | 5.8 (4.6)**              | 10.7 (2.7)d 3–14      |
| Stroop inhibition: scaled score | 6.3 (4.6)**†                 | 3.6 (3.2)**              | 11.5 (2.0)c 7–14      |

a Reading IQ measured on the National Adult Reading Test (NART) (Nelson, 1982) unless the subject scored $<15/50$ on this test, in which case the Schonell Graded Word Reading Test IQ was used (Schonell and Goodacre, 1971).

b Two temporal lobe variant FTLD, one subject with Alzheimer’s disease did not perform synonyms tests.

c Three subjects with temporal lobe variant FTLD, two subjects with Alzheimer’s disease did not perform the London landmarks test.

d 34/35 controls tested on these tasks.

e 19/35 controls tested on these tasks.

f One temporal lobe variant FTLD subject unable to read the words and a scaled score of 1 was used.

g n = 12 (two temporal lobe variant FTLD subjects unable to name colours).

h One temporal lobe variant FTLD subject did not perform recognition memory tasks.

*Significantly worse than controls $(P < 0.01)$; **significantly worse than controls $(P < 0.001)$; †significantly worse than other patient group $(P < 0.05)$; ‡significantly worse than other patient group $(P < 0.001)$.

BPVS = British Picture Vocabulary Scale (McCarthy and Warrington, 1992); Concrete and Abstract Synonyms Test (Warrington et al., 1998); D5 = Wechsler Memory Scale – Revised (WMS-R) digit span (Wechsler, 1987); GDA = Graded Difficulty Arithmetic; GNT = Graded Naming Test (Warrington, 1997); Landmark name/identity = London landmark naming and identification test (Whiteley and Warrington, 1978); ODT = Object Decision Task (Warrington and James, 1991); RMT = Recognition Memory Tests (Warrington, 1984); Stroop, Delis-Kaplan Executive Function System Stroop test (Delis et al. 2001); WASI = Wechsler Abbreviated Scale of Intelligence (Wechsler, 1999).
cues. Vocal samples (each 5 s in duration) were derived from publicly available sources. Twenty-four trials (12 male) were presented; the task was to decide if the voice was male or female.

**Vocal size (vocal tract length)**

Like voice gender, vocal size is determined by lower-level properties: in this subtest we were interested in subjects’ ability to make more fine-grained categorical perceptual judgements of vocal size by manipulating vocal tract length information in isolation, as previously described in normal subjects (Ives et al., 2005). Stimuli were based on 10 consonant–vowel syllables recorded by a single male speaker and digitally resynthesized using a previously described algorithm (Kawahara and Irino, 2004) that allows apparent vocal tract length to be varied independently of glottal pulse rate (voice pitch). Each syllable was presented at two extreme vocal tract length values, one corresponding to a speaker height of 2 m (equivalent to a very tall male, ‘big’) and the other to a height of 0.5 m (equivalent to a child, ‘small’), and randomly assigned one of four pitches within the normal human male vocal range (116, 120, 172, 190 Hz), which was varied independently of vocal tract length. Examples of stimuli are available from the authors. Twenty trials (10 ‘big’, 10 ‘small’) were presented; on each trial, subjects heard a sequence of four repetitions of the same stimulus, and were asked to judge if the sounds were made by a big person or a small person.

**Experimental behavioural investigations: voice recognition**

In this test we aimed to assess different aspects of semantic processing of famous voices (voice recognition) comprising familiarity, naming, identification from biographical information and cross-modal matching, in comparison to recognition of faces and names for the same famous individuals. Sixty voice samples and face photographs were initially obtained from publicly available sources on the internet and chosen so as to minimize other potential semantic cues to recognition. A set of 24 voice samples of well-known public figures was selected from this larger set, based on a pilot analysis in a separate group of healthy older British controls (Hailstone et al., 2010); the best-recognized samples were included in the final stimulus set used in all semantic subtests. These 24 famous individuals (Supplementary Table 1) comprised 10 politicians, five actors, seven other media personalities from television and radio, and two members of the British royal family. Examples of stimuli are available from the authors. Face photographs and names of the same set of 24 famous individuals were used for the face and name processing tasks, respectively.

**Familiarity of voices, faces and names**

In this subtest the set of 24 famous voices was supplemented by 24 unfamiliar voices and faces (as classified by >75% of healthy controls in the pilot study: Hailstone et al., 2010) matched by gender to the famous set and approximately matched for age and accent. The written names of the same 24 famous individuals were supplemented with 24 fabricated personal name foils. For each modality, 48 trials (24 famous, 24 unfamiliar) were presented; each stimulus was presented once, and the task was to decide if the stimulus was familiar in a forced choice (‘yes – no’) protocol. As voice familiarity was the primary focus here, voices were presented first (in order to minimize priming effects in the voice modality), followed by faces and names.

**Identification of voices, faces and names**

In this subtest, the task was to identify the person as precisely as possible; for voices and faces, if the subject was not able to name the person, they were asked to provide other biographical details (e.g. an event closely associated with the person, occupational information), in line with the criteria used by Snowden et al. (2004). For names, on each trial the subject was required to provide identifying information about the person. For voices, national or regional origin was not accepted as evidence of person recognition, since this could be based on accent cues alone. As voice identification was the focus here, voices were presented first (in order to minimize priming effects in the voice modality), followed by faces and names.

**Cross-modal recognition of voices and faces**

We employed a cross-modal matching task in order to allow patients to demonstrate recognition of voices and faces using an alternative procedure that did not rely on word retrieval. For both face and voice targets, three stimulus arrays were selected using individuals from the set of 24 individuals; each individual was represented in one of the arrays. The first array contained the six females from the complete set, a second array contained the nine male politicians, and the third contained the nine male media figures (as career is likely to be an important organizational principle in the domain of person knowledge; Crutch and Warrington, 2004). The set of 24 faces was presented first, and the task was to match the face to one of the
names in the array. The set of 24 famous voices was presented with the same arrays but with simultaneous presentations of faces and names in each array; the task was to match the voice to one of the face-name pairs.

Experimental behavioural investigations: general procedure

The experimental tests were administered to subjects over several sessions. Auditory stimuli were presented from digital wavefiles on a laptop computer via headphones at a comfortable listening level in a quiet room. Visual stimuli were presented as high-quality greyscale photographs. Verbal stimuli were simultaneously presented as written words and spoken by the examiner (control subjects were presented with written words only). All stimuli were presented in fixed randomized order. Subject responses were collected for off-line analysis. Before beginning each test, several practice trials were administered to ensure the subject understood the task. No feedback about performance was given during the test. Voice stimuli were presented once only for all tasks, with the exception of the famous voice identification task, where subjects were permitted a further presentation if requested. No time limit was imposed.

Experimental behavioural investigations: analyses

Experimental behavioural data were analysed in STATA R9.2 (Stata Corporation). For the perceptual tests, differences in mean scores between groups were assessed using z-tests and 95% Wald-type confidence intervals, with standard errors calculated using bootstrapping (2000 replicates). For the semantic subtests, the effect of stimulus presentation modality was assessed using a bootstrapped linear regression model with 2000 replicates, which allowed for the repeated measures from subjects. A global Wald test of interaction was carried out to test the hypothesis that group differences in scores varied between modalities, and modality-associated differences in performance between the temporal variant FTLD and Alzheimer’s disease groups were assessed in pair-wise comparisons between modalities. Using this model, differences between the two patient groups were adjusted for modality performance differences exhibited by healthy controls (anticipated from previous work; Hailstone et al., 2010).

Within each patient group, correlation coefficients between experimental tests were estimated with 95% bias-corrected bootstrap confidence intervals (2000 replicates). Correlations were estimated between perceptual discrimination subtest scores; between semantic subtest scores within and between modalities; and between perceptual and semantic performance. Associations with disease severity measures were also assessed, using linear regression models with 95% Wald-type bootstrap confidence intervals with 2000 replicates. As severity measures, symptom (clinical disease) duration was used for both disease groups; in addition, Mini-Mental State Examination score was used for off-line analysis. Before beginning each test, several practice trials were administered to ensure the subject understood the task. No feedback about performance was given during the test. Voice stimuli were presented once only for all tasks, with the exception of the famous voice identification task, where subjects were permitted a further presentation if requested. No time limit was imposed.

Neuroimaging protocol and analysis

Brain image acquisition

For 18 patients with Alzheimer’s disease and 11 patients with temporal lobe variant FTLD, T1-weighted volumetric magnetic resonance images were acquired on a Siemens Trio TIM 3T scanner (Siemens Medical Systems). Scans were acquired using a 3D magnetization prepared rapid gradient echo (MP-RAGE) sequence producing 208 contiguous 1.1 mm thick sagittal slices with 28 cm field of view and a 256 × 256 acquisition matrix, giving approximately isotropic 1.1 mm cubic voxels.

Voxel-based morphometry analyses

Magnetic resonance brain images were processed using MATLAB 7.2 (The MathWorks Inc.) and SPM8 software (Statistical Parametric Mapping, Version 8; http://www.fil.ion.ucl.ac.uk/spm) with default settings for all parameters; normalization was performed using the DARTEL toolbox (Ashburner, 2007). Further details of preprocessing steps are provided in the Supplementary material.

For each modality in the experimental battery (voices, faces, names), associations between regional grey matter volume and subtest performance were assessed in both disease groups using linear regression models. In separate-modality design matrices, grey matter volume was modelled as a function of the experimental subtest score-by-group interaction term with group, age and total intracranial volume. The latter was measured outside statistical parametric mapping using a previously described procedure (Whitwell et al., 2001). In addition to these separate-modality analyses, joint combined-modalities models were used to assess the independent partial associations of voice, face and name modalities for the familiarity subtest and the identification subtest and partial associations of voice and face modalities for the cross-modal recognition subtest. For each subtest, F tests were used to assess grey matter associations with performance for each modality (adjusting for the others) and conjointly across modalities.

An explicit analysis mask was used to exclude any voxels for which >20% of the images had an intensity value of <0.1 (Ridgway et al., 2009). Grey matter associations were assessed over the whole brain and within the regions of interest specified by our prior anatomical hypotheses; small volumes covering temporal and parietal lobe regions specified in our prior anatomical hypotheses were created manually in MRICron® (http://www.cabiatl.com/micro/mricron/index.html) from a study-specific template image (further details in Supplementary material). A voxel-wise statistical threshold P < 0.05 family-wise-error-corrected for multiple comparisons was applied in all analyses (a global P < 0.05 family-wise error-corrected threshold was applied in the combined-modalities conjunction analysis). Statistical parametric maps were displayed as overlays on the study-specific template. A voxel-wise exclusive masking procedure was applied to display grey matter areas associated with voice processing performance but not with performance in other modalities. In order to report coordinates of local maxima in the standard stereotactic Montreal Neurological Institute space, the grey matter segment of the final DARTEL template was affine registered to the a priori grey matter tissue probability map in statistical parametric maps, and the DARTEL coordinates were transformed using the estimated affine mapping to Montreal Neurological Institute space.
Results

General neuropsychological data

Results of the general neuropsychological assessment are presented in Table 2. Relative to the healthy control group both the temporal lobe variant FTLD group and the Alzheimer’s disease group showed reduced verbal and performance IQ and deficits of executive function and cognitive speed, recognition memory for words and faces, naming and calculation; the Alzheimer’s disease group showed additional deficits of visual object perception and auditory verbal working memory. The temporal lobe variant FTLD group had lower verbal and reading IQ than the Alzheimer’s disease group and performed significantly worse than the Alzheimer’s disease group on tests of naming, tests of semantic knowledge (verbal comprehension and London landmark identification tests), reading and recognition memory for faces; while the Alzheimer’s disease group performed significantly worse than the temporal lobe variant FTLD group on tests of non-verbal reasoning and recognition memory for words. Considered together, these patterns of performance support the clinical and neuroanatomical classification for each disease group.

Peripheral hearing data

On screening assessment of peripheral hearing, increasing age was associated (as anticipated) with a significant increase in mean detection threshold at the three highest frequencies tested (2, 3 and 4 kHz). Relative to the healthy control group, there was a significant difference (P < 0.05) in mean detection thresholds at 0.5 kHz for both patient groups (adjusted differences in means from controls: Temporal lobe variant FTLD group = 7.2 dB, Alzheimer’s disease group = 4.7 dB); these threshold elevations were small and unlikely to be clinically relevant. No significant differences were shown by either patient group with respect to controls at any other frequency tested, and no significant differences were observed between temporal lobe variant FTLD and Alzheimer’s disease groups at any frequency.

Experimental behavioural data

Here we report the findings from the group analyses on tests in the experimental battery. Details of individual subject data are provided in the Supplementary material.

Perceptual processing of voice attributes

Results for the patient and healthy control groups on early perceptual and apperceptive subtests for each modality are summarized in Table 3 and further details are provided in Supplementary Table 2.

On the vocal gender subtest, the Alzheimer’s disease group performed significantly worse (P < 0.05) than the healthy control group, however this difference was driven by a subgroup of four patients with Alzheimer’s disease (the remaining patients scored at ceiling on this task). The performance of the temporal lobe variant FTLD group did not differ from healthy controls, however all subjects in the temporal lobe variant FTLD and control groups performed at ceiling on this subtest. On the vocal size subtest, there were no significant group performance differences and a large range of scores in all three groups.

On both the ‘easy’ and the ‘difficult’ speaker discrimination subtests, the Alzheimer’s disease group performed significantly worse (P < 0.05) than healthy controls. There were no significant performance differences between the temporal lobe variant FTLD group and healthy controls or between the two patient groups.

Recognition of voices, faces and names

Results for the patient and healthy control groups on semantic subtests for each modality are summarized in Table 4 and further details are provided in Supplementary Tables 2, 3 and 4.

On all semantic subtests, both the temporal lobe variant FTLD group (P < 0.001) and the Alzheimer’s disease group (P < 0.05)

| Subtest (max score)          | Temporal lobe variant FTLD n = 14 | Alzheimer’s disease n = 22 | Healthy controls n = 35 | Temporal lobe variant FTLD–Alzheimer’s disease | Difference in means (95% confidence interval) |
|-----------------------------|-----------------------------------|-----------------------------|-------------------------|-----------------------------------------------|-----------------------------------------------|
| Voice perception            |                                    |                             |                         |                                               |                                               |
| Size perception (/20)        | 16.7 (2.8) 11–20                   | 17.4 (2.1) 12–20            | 17.1 (2.9) 9–20         | –0.7 (–2.4, 1.0)                              |                                               |
| Gender perception (/24)      | 24.0 (0.0) 24–24                   | 23.7 (0.6)* 22–24           | 24 (0.0) 24–24          | 0.3 (0.01, 0.5)*                              |                                               |
| Easy speaker discrimination  | 24.7 (1.6) 22–27                   | 24.1 (3.2)* 15–28           | 25.6 (1.5) 21–28        | 0.6 (–1.0, 2.2)                               |                                               |
| Difficult speaker discrimination (/12) | 9.2 (1.2) 7–11   | 8.8 (1.7)** 6–12           | 9.9 (1.4) 7–12          | 0.4 (–0.5, 1.4)                               |                                               |
| Face perception              |                                    |                             |                         |                                               |                                               |
| Benton facial recognition test (/56) | 42.8 (4.0)** 37–50 | 42.2 (5.8)** 32–52 | 48.0 (3.2) 42–56       | 0.7 (–2.5, 3.8)                               |                                               |

*Significantly worse than controls (P < 0.05); **significantly worse than controls (P < 0.01); †Alzheimer’s disease group significantly worse than temporal lobe variant FTLD group (P < 0.05).
performed significantly worse than the healthy control group. For both disease groups and also for the healthy control group, mean absolute scores across semantic subtests were lower for voice recognition than for recognition in the other modalities. The temporal lobe variant FTLD group performed significantly worse than the Alzheimer’s disease group ($P < 0.01$) on all familiarity subtests apart from face familiarity [for which there was a trend ($P = 0.07$) to worse performance], on all identification subtests in each modality, on the cross-modal subtests and on voice and face naming.

There was a significant interaction between group and modality for all subtests: familiarity ($P < 0.001$), identification ($P < 0.05$), cross-modal recognition ($P < 0.01$) and naming ($P < 0.01$). The temporal lobe variant FTLD group showed a significantly larger ($P < 0.05$) decrease in score compared to the Alzheimer’s disease group for identification in the name modality compared with recognition in the other modalities. The temporal lobe variant FTLD–Alzheimer’s disease performance discrepancy did not differ significantly between modalities for the other subtests. In general, significant performance correlations were observed between voice recognition subtests in the Alzheimer’s disease group though less consistently in the temporal lobe variant FTLD group; while voice and face identification performance was correlated in both disease groups (Supplementary Table 3). Voice identification performance was not closely correlated with apperceptive performance in either disease group (Supplementary Table 2). Voice identification and familiarity performance were not correlated with disease severity measures in either disease group (Supplementary Table 4).

Right versus left temporal lobe damage in temporal lobe variant FTLD

Temporal lobe variant FTLD subgroups with predominantly left-sided ($n = 9$) versus predominantly right-sided ($n = 4$) temporal lobe atrophy did not show significant performance differences on any of the voice processing subtests. Further details are provided in the Supplementary material.

Neuroanatomical data

Interactions between disease group and performance

No significant grey matter associations were identified for group–performance interactions for any of the experimental subtests over the whole brain volume. Restricting analyses to the pre-specified anatomical volume of interest there was a significant interaction between group and performance on the ‘easy’ speaker discrimination task in the right parahippocampal gyrus (local maximum Montreal Neurological Institute coordinates: 35-51-6; cluster size 123 voxels, $P < 0.05$ after family-wise error correction). Voice discrimination performance in the Alzheimer’s disease group (but not the temporal lobe variant FTLD group) was positively associated with grey matter in right inferior parietal cortex ($P < 0.05$ after family-wise error correction over the pre-specified small volume of interest; Table 5); additional associations of voice discrimination were present in right parahippocampal gyrus and left inferior parietal cortex at an uncorrected threshold ($P < 0.001$ over the whole brain volume; Fig. 1).
Associations of performance across disease groups

The results of the neuroanatomical analysis across both the temporal lobe variant FTLD and Alzheimer’s disease groups (adjusting for group membership) are summarized in Table 5; statistical parametric maps are presented in Fig. 1.

We consider first the results of analyses for associations of experimental test performance over the whole brain volume. No significant associations of voice perceptual performance across both disease groups were identified. In the separate-modality analyses of semantic processing, cross-modal recognition of voices and identification and cross-modal recognition of faces were each positively associated with grey matter volume at the right temporal pole; in addition, cross-modal recognition of voices and identification of faces and names were each positively associated with grey matter volume in right anterior fusiform gyrus (all $P < 0.05$ after family-wise error correction over the whole brain volume). In the combined-modalities analysis, there was a common grey matter association of voice, face and name identification at the right temporal pole ($P < 0.05$ after family-wise error correction over the whole brain volume), however no significant partial associations of voice, face or name identification were identified.

Restricting analyses to the pre-specified anatomical volumes of interest, a number of additional associations were identified (all $P < 0.05$ after family-wise error correction over the relevant small volume). In the separate-modality analyses of semantic processing, across both disease groups, voice and name identification were each positively associated with grey matter at the right temporal pole; voice identification (but not face or name identification) was positively associated with grey matter in right amygdala and hippocampus, while face and name identification (but not voice identification) were each positively associated with grey matter at the left temporal pole. In the combined-modalities analysis, a common grey matter association of voice, face and name familiarity was identified in right fusiform gyrus; common grey matter associations of voice and face cross-modal recognition were identified in right temporal pole and anterior fusiform.

### Table 5 Voxel-based morphometry data: neuroanatomical associations of behavioural performance

| Task | Neuroanatomical associations | Cluster size (voxels) | Local maxima Coordinates (mm) | Z-score |
|------|------------------------------|-----------------------|-------------------------------|---------|
|      | Side | Area                                   |                             |         |
|      |      |                                        | 56                           | 42      |
| Apperceptive | Voice discrimination* | Inferior parietal cortex | 838                           |         |
|      | L    | Temporal pole                           | 678                          |         |
|      | R    | Anterior middle temporal gyrus          | 250                          |         |
|      | R    | Anterior fusiform gyrus                 | 1203                         |         |
|      |      |                                        | 28                           | 42      |
|      |      |                                        | 28                           | 42      |
|      |      |                                        | 35                           | 22      |
|      |      |                                        | 35                           | 22      |
|      | Face |                                        | 3829                         |         |
|      | R    | Temporal pole                           | 558                          |         |
|      | L    | Temporal pole                           | 481                          |         |
|      | R    | Anterior fusiform gyrus*                | 18                           |         |
|      | R    | Anterior fusiform gyrus*                | 18                           |         |
|      | L    | Temporal pole                           | 942                          |         |
|      | R    | Temporal pole*                         | 1861                         |         |
|      |      |                                        | 22                           | 42      |
|      |      |                                        | 22                           | 42      |
|      |      |                                        | 22                           | 42      |
|      |      |                                        | 22                           | 42      |
|      |      |                                        | 22                           | 42      |
|     |      |                                        | 22                           | 42      |
| Cross-modal matching | Voice | Temporal pole*                          | 16                           |         |
|      | R    | Anterior fusiform gyrus*                | 3098                         |         |
|      | R    | Anterior fusiform gyrus*                | 2712                         |         |
|      | R    | Anterior fusiform gyrus                 | 2712                         |         |
|      |      |                                        | 25                           | 46      |
|      |      |                                        | 25                           | 46      |
|      |      |                                        | 25                           | 46      |

Results for voice discrimination were derived from the Alzheimer’s disease group only; all other results were derived across the temporal lobe variant FTLD and Alzheimer’s disease groups. All clusters of size $>10$ voxels are presented.

* ‘Easy’ version of speaker discrimination task (see text).

* Results based on combined-modalities analyses; other results based on separate-modality analyses (see text).

* Areas with local maxima exceeding voxel-wise significance threshold $P < 0.05$ after family-wise error correction over the whole brain; other local maxima after correction over the pre-specified small volume of interest (co-ordinates in Montreal Neurological Institute stereotactic space).
gyrus. No significant partial associations were identified for voice, face or name familiarity or for cross-modal recognition of voices or faces. No significant grey matter associations of voice or face naming performance were identified.

**Discussion**

Here we have described behavioural and neuroanatomical signatures of voice processing deficits in FTLD and Alzheimer’s disease. The key findings from the study are two-fold. In behavioural terms, impairments of voice recognition are significant in these canonical dementia syndromes but particularly severe in temporal lobe variant FTLD. In neuroanatomical terms, impaired voice recognition is associated with grey matter loss in the right anterior temporal lobe, suggesting that this region is critical for voice recognition as well as other modalities of person knowledge.

On all behavioural measures of voice recognition (familiarity, identification, naming and cross-modal identity matching), both disease groups showed deficits relative to healthy controls and the temporal lobe variant FTLD group performed significantly worse than the Alzheimer’s disease group. Despite substantial individual variation in performance, both disease groups showed a clear overall trend to conjoined deficits of voice recognition with deficits of other modalities of person knowledge (in particular, recognition of faces). The largest performance discrepancy between the disease groups occurred for the processing of personal names. On measures of perceptual processing of voices, the Alzheimer’s disease group (but not the temporal lobe variant FTLD group) showed deficits relative to healthy controls, suggesting that impairments of voice perception may show relative specificity for Alzheimer’s disease.

The neuroanatomical analysis showed that recognition measures for voice, face and name processing were associated across both disease groups with grey matter volume in right temporal pole and anterior fusiform gyrus. Similar regions have been implicated in the processing of familiar voices by the healthy brain (Belin et al., 2000, 2002; Nakamura et al., 2001; Shah et al., 2001; Belin and Zatorre, 2003; von Kriegstein et al., 2003, 2005; von Kriegstein and Giraud, 2004; Andics et al., 2010). These same areas are sites of heavy disease involvement in temporal lobe variant FTLD, consistent with the more severe person recognition deficits in this group compared with the Alzheimer’s disease group. However, it is unlikely the profile of anatomical associations observed was driven entirely by the temporal lobe variant FTLD group, since there was no evidence of significantly different grey matter associations of semantic test performance between the disease groups. We do not wish to over-emphasize this apparent anatomical convergence. Whereas Alzheimer’s disease is pathologically homogeneous, temporal lobe variant FTLD is likely to
be pathologically heterogeneous (Josephs et al., 2011): a shared macro-anatomical substrate may be underpinned by distinct patterns of cellular involvement and correspondingly distinct pathophysiological mechanisms. Moreover, it remains possible that there is a more fine-grained segregation of processing for different modalities within the relatively large cortical areas identified here using voxel-based morphometry (Olson et al., 2004).

Certain mesial temporal lobe structures (amygdala, hippocampus and entorhinal cortex) showed an association with voice identification but not other modalities of person knowledge at the prescribed threshold (Fig. 1): the mesial temporal lobe has been previously implicated in familiar voice processing by healthy subjects (Nakamura et al., 2001; von Kriegstein and Giraud, 2004; von Kriegstein et al., 2005; Andics et al., 2010) and this region may be involved in processing sensory object information (Lee et al., 2005) and particularly in tracking information in sound (for example, familiar musical melodies) (Samson and Zatorre, 2008). However, any apparent modality-specificity here should be interpreted with caution; no independent associations of recognition in a particular modality emerged when modalities were assessed together.

Both the behavioural and neuroanatomical findings here are consistent with a growing body of evidence implicating the anterior temporal lobe in multi-modal processing of semantic knowledge and more particularly, person knowledge (Bozeat et al., 2000; Coccia et al., 2004; Luzzi et al., 2007; Rami et al., 2007; Lambon Ralph and Patterson, 2008). Previous functional imaging and lesion evidence suggests that the temporal pole and anterior fusiform participate in a cooperative network mediating different aspects of semantic processing (Ellis et al., 1989; Tranel et al., 1997; Papagno and Capitani, 1998; Grabowski et al., 2001; Thompson et al., 2004; von Kriegstein et al., 2005; Tranel, 2006; Mion et al., 2010). While we cannot specify the precise role of anterior fusiform and temporal pole in the present study, grey matter volume in anterior fusiform was associated with performance on familiarity and cross-modal judgements across modalities, consistent with the operation of person identity nodes in this area (Belin et al., 2004). We do not argue that processing voice familiarity is a purely semantic capacity; analogously with familiarity for other kinds of stimuli, voice familiarity may also have perceptual, affective and executive dimensions (Gainotti, 2007; Lucchelli and Spinnler, 2008). Taken together, our findings support the existence of multi-modal deficits of person knowledge in FTLD and Alzheimer’s disease. Rather than a purely amodal or fully multi-modal organization, the data suggest that verbal (name) and non-verbal (voice, face) modalities of person knowledge may be partially differentiated, whereas modalities of non-verbal person knowledge are more closely aligned (Warrington, 1979; Snowden et al., 2004). The present data (based on small case numbers) do not resolve the important issue of potentially asymmetric temporal lobe contributions to different components of person knowledge (Snowden et al., 2004).

In this study, deficits of voice perception were restricted to the Alzheimer’s disease group, and involved voice apperception (speaker discrimination) and encoding of one perceptual attribute (vocal gender) while sparing encoding of another attribute (vocal size). A neuroanatomical association of apperceptive performance in the Alzheimer’s disease group was identified in the right inferior parietal lobe. The present findings underline the potential for development of semantic deficits of voice recognition (and other aspects of person recognition) despite intact pre-semantic perceptual mechanisms; however, deficits of voice perception may have contributed to the development of voice recognition impairment in the Alzheimer’s disease group. Functional imaging work in healthy subjects has delineated a network of cortical areas including the parietal lobe for processing voice information under non-canonical listening conditions (Bishop and Miller, 2009). We propose that inferior parietal cortex is involved in the structural representation of voices (Bruce and Young, 1986; Burton et al., 1990; Belin et al., 2004), perhaps by holding voice information online in working memory for comparison with incoming alternative auditory ‘views’ of the speaker (e.g. the same voice speaking different phonemes).

This study suggests clear directions for future work. It has been proposed that modality-specific deficits of person knowledge become generalized with the evolution of neurodegenerative disease (Evans et al., 1995; Gentileschi et al., 1999; Gainotti et al., 2003, 2008); the present study suggests that the profile of development of deficits may hold information about the organization of processing within and between modalities. This issue will only be addressed by longitudinal studies based on a systematic analysis of different levels of processing and comparing modalities and disease groups. Whereas semantic processing of voices is relatively easily investigated by adapting standard neuropsychological techniques, a detailed understanding of voice perception and its disorders will require the design of customized stimuli that allow particular vocal attributes to be isolated and manipulated. Finally, there is a need for correlation of voice processing measures with structural and functional anatomical data and with tissue histopathology in a range of neurodegenerative diseases.

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Supplementary material

Supplementary material is available at Brain online.

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