Music models aberrant rule decoding and reward valuation in dementia

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Abstract

Aberrant rule- and reward-based processes underpin abnormalities of socio-emotional behaviour in major dementias. However, these processes remain poorly characterized. Here we used music to probe rule decoding and reward valuation in patients with frontotemporal dementia (FTD) syndromes and Alzheimer’s disease (AD) relative to healthy age-matched individuals. We created short melodies that were either harmonically resolved (‘finished’) or unresolved (‘unfinished’); the task was to classify each melody as finished or unresolved (rule processing) and rate its subjective pleasantness (reward valuation). Results were adjusted for elementary pitch and executive processing; neuroanatomical correlates were assessed using voxel-based morphometry. Relative to healthy older controls, patients with behavioural variant FTD showed impairments of both musical rule decoding and reward valuation, while patients with semantic dementia showed impaired reward valuation but intact rule decoding, patients with AD showed impaired rule decoding but intact reward valuation and patients with progressive non-fluent aphasia performed comparably to healthy controls. Grey matter associations with task performance were identified in anterior temporal, medial and lateral orbitofrontal cortices, previously implicated in computing diverse biological and non-biological rules and rewards. The processing of musical rules and reward distills cognitive and neuroanatomical mechanisms relevant to complex socio-emotional dysfunction in major dementias.

Key words: music; reward; emotion; Alzheimer’s disease; frontotemporal dementia

Introduction

Disturbances of complex emotional and social behaviour are a defining hallmark of frontotemporal dementia (FTD) and may be prominent in other neurodegenerative diseases, notably Alzheimer’s disease (AD) (Rascovsky et al., 2011; Sturm et al., 2013; Warren et al., 2013; Sapey-Triomphe et al., 2015; Clark et al., 2016). The spectrum of socio-emotional dysfunction in these diseases is complex and poorly understood; however, certain core themes emerge. These include impaired understanding of social norms and their violation (potentially leading to phenomena such as faux pas, loss of decorum, insensitivity to others’ discomfiture and diffusion of social boundaries: Ibanez and Manes, 2012) and abnormal affective responses to homeostatic...
signals (leading, e.g. to apathy or abnormal hedonic investment in potentially rewarding, aversive or banal stimuli: Zhou and Seeley, 2014), as well as altered reactivity to emotional cues from complex environments (Sturm et al., 2015) or reduced ability to use semantic regularities in anticipating future events (Bertoux et al., 2014; Irish and Piolino, 2016). These processes might be collectively conceptualized as impaired ‘rule’ decoding and altered reward valuation from socio-emotional and other kinds of signals. Behaviourally, a major unifying goal of such processes is to respond appropriately to salient events in the world at large: ‘salience’ is a property often associated with stimuli that violate expectations (‘rules’) established by prior experience of related stimuli and salient stimuli frequently have reward or aversive potential, according to their intrinsic (biological or cognitive) value and behavioural context (Wallis, 2007; Lang et al., 2009; Dalton et al., 2012; Ibanez and Manes, 2012; Schultz, 2013; Zhou and Seeley, 2014).

Emerging evidence in neurodegenerative diseases has underlined the clinical and neuroanatomical relevance of rule and reward processing. Large-scale cerebral networks coding salience (comprising insula, anterior cingulate and orbitofrontal cortex) and reward value (comprising ventral striatal and mesolimbic dopaminergic circuitry) have been implicated in the pathogenesis of socio-emotional phenotypes in FTD (Wallis, 2007; Lang et al., 2009; Fievani et al., 2011; Halabi et al., 2013; Perry et al., 2014, 2015; Zhou and Seeley, 2014; Bocchetta et al., 2016). Altered interactions between salience, reward and temporo-parietal ‘default-mode’ networks underpin socio-emotional dysfunction in AD (Ismaiel et al., 2008; O’Reilly et al., 2013; Zhou and Seeley, 2014; Fletcher et al., 2015b; Perry et al., 2015; Ahmed et al., 2016). Deficient integration of signal processing with central reward evaluation and autonomic regulatory mechanisms may constitute a common effector pathway in a range of diseases (Zhou and Seeley, 2014; Fletcher et al., 2015a,b). However, the study of rule processing, salience detection and associated reward processing in the dementias poses a number of challenges. These include the complexity of natural socio-emotional scenarios, the diversity of contingent phenomena (spanning the gamut of primary biological and secondary hedonic stimuli) and the difficulty of manipulating relevant stimulus parameters under experimental conditions.

As a candidate model system for analysing rule decoding and reward valuation in dementia, music has a number of attractive attributes. Music is universal, ubiquitous and pleasurable for most listeners; its effects are grounded in cerebral circuitry that mediates pattern resolution and hedonic processing, overlying the distributed networks implicated in FTD and AD (Koelsch et al., 2000; Blood and Zatorre, 2001; Janata et al., 2002; Steinbeis et al., 2006; Pressnitzer et al., 2011; Salimpoor et al., 2011, 2013, 2015; Seger et al., 2013; Clark et al., 2014, 2015a). The socio-emotional resonance of music suggests it may be a fertile model for exploring the building blocks of more complex interpersonal and affective behaviours in daily life (Clark et al., 2015a). Music necessarily unfolds over time, mirroring the dynamic nature of socio-emotional interactions. On the other hand, despite its propensity to engage biological reward circuitry, music is an autonomous signalling system that does not depend on extra-musical associations. Moreover, the characteristics of music that convey salience and hedonic value are highly codified and relatively straightforward to vary experimentally. The ‘rules’ governing harmonic structures are acquired implicitly by normal listeners through exposure to the music of the dominant culture and are engaged rapidly and automatically even during non-attentive listening (Tillmann, 2005; Huron, 2006; Loui et al., 2009; Koelsch et al., 2013; Choi et al., 2014). Indeed, music may be an ideal probe of the interface between cognition and emotion, via the medium of psychological expectancy (Krumhansl, 1997; Huron, 2006).

The relations between rule-based musical expectancy, arousal and valence (perceived pleasantness) are complex and influenced by the specific musical context. The interplay of tension and resolution is a fundamental mechanism underlying a range of psychophysiological (arousal and valence) effects associated with the manipulation of musical structure (Krumhansl, 1997; Steinbeis et al., 2006; Gingras et al., 2016; Tsai and Chen, 2015). Musical pleasure and reward (though separable phenomena) are typically closely correlated (Salimpoor et al., 2013): although ambiguity or surprise in more complex music and musical ‘scenes’ can be intensely pleasurable (Huron, 2006; Pressnitzer et al., 2011), melodies with a harmonic structure that fulfils expectation or resolves ambiguity tend to reduce subjective tension and are usually perceived as subjectively pleasurable or rewarding, while lack of resolution or confounded expectation is associated with increased subjective tension and negative affect (Steinbeis et al., 2006, Gingras et al., 2016; Tsai and Chen, 2015; Tsai et al., 2015). This propensity of music to create psychological expectancy (‘pay-off’ vs disappointment) links it naturally to reward valuation, and indeed this is already recognized in the music neuroscience literature (Li et al., 2015; Tsai and Chen, 2015). Reward is a complex, multidimensional psychophysiological construct, but generally entails the potential for consummatory behaviour or active updating of stimulus value, completion or resolution even in the absence of overt goal-directed action (Schultz, 2013; Perry et al., 2015).

From a clinical perspective, altered behavioural responses to music are well documented in many patients with dementia and may stratify FTD and AD syndromes. Recent evidence suggests that behavioural variant (bv)FTD may impair the analysis of musical structure (in particular, anticipation of structure unfolding in time) as well as hedonic valuation of music, whereas semantic dementia (SD) may be associated with relatively preserved processing of musical structure and progressive non-fluent aphasia (PNFA) and AD with relatively preserved hedonic valuation of music (Omar et al., 2010; Weinstein et al., 2011; Haieh et al., 2012; Downey et al., 2013; Fletcher et al., 2013, 2014, 2015b; Henley et al., 2014; Agustus et al., 2015).

Here we used music as a paradigm to assess rule decoding and reward valuation in the canonical syndromes of FTD and AD, referenced to healthy older individuals. We presented novel melodies that either resolved according to the rules of classical harmony or did not resolve: the behavioural task was to decide whether or not each melody resolved (rule decoding) and to rate its pleasantness (reward valuation). We adopted this tonal expectancy task (rather than simply assessing, e.g. detection of dissonant or scale-deviant notes) because we wished to model the dynamic scenarios of natural, extra-musical, socio-emotional behaviour in daily life: in more complex social scenarios, the potential for reward (here, modelled as harmonic resolution) rather than the delivery of ‘punishment’ (a dissonant note) tends to be the prime mover of behaviour (Milinski and Rockenbach, 2012). At the same time, we wished to avoid any requirement for explicit labelling of musical emotions, a somewhat artificial task that is known to be impaired in FTD syndromes (Omar et al., 2011). We assessed neural correlates of musical rule and reward processing using voxel-based morphometry of patients’ brain MR images. Based on previous evidence, we hypothesized that bvFTD would be associated with abnormalities of both musical rule decoding (categorization of
tonally resolved vs unresolved melodies) and reward valuation (emotional rating of these melodies) (Omar et al., 2010; Downey et al., 2013; Henley et al., 2014; Agustus et al., 2015; Fletcher et al., 2015b) while SD would show abnormal musical reward valuation despite relatively preserved rule decoding (Omar et al., 2010; Weinstein et al., 2011; Hsieh et al., 2012; Fletcher et al., 2013, 2015b) and PNFA and AD would show abnormal musical rule decoding despite relatively preserved reward valuation (Rohrer et al., 2012; Fletcher et al., 2015b; Golden et al., 2017). Based on previous work in the healthy brain and in patients with focal brain damage, we further hypothesized that harmonic classification of melodies would have a neuroanatomical correlate in superior temporal and prefrontal areas that mediate the analysis of musical and other rule-based patterns (Koelsch et al., 2000, 2013; Janata et al., 2002; Salimpoor et al., 2013, 2015; Seger et al., 2013; Clark et al., 2015a); while the rating of harmonic pleasantness would have a correlate in inferior frontal areas (including orbitofrontal cortex and inferior frontal gyrus) implicated in the affective labelling of music and other stimuli (Blood and Zatorre, 2001; Salimpoor et al., 2011, 2013, 2015; Clark et al., 2014, 2015a) and the integration of semantic and emotional information (Koelsch et al., 2006; Belyk et al., 2017).

Materials and methods

Participants

In total 14 patients with typical AD (henceforth, ‘AD’) led by episodic memory decline, 11 patients with bvFTD, 6 patients with SD and 8 patients with PNFA were recruited. All patients fulfilled consensus clinical criteria for the relevant syndromic diagnosis (Gorno-Tempini et al., 2011; Rascovsky et al., 2011; Dubois et al., 2014). Twenty-two healthy older individuals with no history of neurological or psychiatric disorders also participated. Demographic details (including an assessment of past musical training and current listening habits) and clinical characteristics of the study cohort are summarized in Table 1. None of the participants had a history of clinically significant hearing loss or congenital amusia. One patient in the bvFTD group and five patients in the SD group had a history of abnormal craving for music or ‘musciophilia’ (Fletcher et al., 2013). All participants had a comprehensive general neuropsychological assessment (summarized in Table 1) and audiometric screening of peripheral hearing function (adapted from a commercial screening audiometry package; details in Supplementary Material). Brain imaging (MRI/CT) revealed a profile of atrophy compatible with the syndromic diagnosis in all patients; no brain images showed a significant cerebrovascular burden. In total 10 of 11 patients in the AD group for whom CSF was available had a protein marker profile suggesting underlying AD pathology (total CSF tau: beta-amyloid, 1 -42 ratio >1, based on local laboratory reference ranges); CSF findings in 11 of 13 patients with other syndromes provided no evidence for underlying AD pathology. At the time of assessment, 13 patients in the AD group were receiving symptomatic treatment with donepezil and 2 with memantine; 1 other patient (in the PNFA group) was receiving donepezil.

The study was approved by the local institutional ethics committee and all participants gave informed consent in accordance with Declaration of Helsinki guidelines.

Musical assessment

Assessment of tonal expectancy. To assess processing of tonal expectancy (musical harmonic) rules and reward value, short monophonic melodies were composed by an experienced musician (O.M.). Melody endings were manipulated such that the melody sounded ‘finished’ (tonally resolved) or ‘unfinished’ (tonally unresolved). Melodies in the ‘finished’ and ‘unfinished’ conditions were closely matched for loudness, length, timbre, instrumentation, key, pitch velocity and tempo. All note sequences were synthesized with piano timbre as digital wave-files using Logic Pro X. Tempo was fixed at 120 beats/min for all stimuli; melodies varied in length between three and five bars; however, the two conditions did not differ in mean length (number of bars or duration; see Supplementary Material). Examples of stimuli are presented in Figure 1; the complete set is notated in Supplementary Figure S1, together with further details of stimulus preparation and pilot testing.

Melodies were presented in randomized order and the task on each trial was to decide first, if the melody sounded ‘finished’ or ‘unfinished’; and second, to rate how pleasing was the ending of the melody (‘How did the tune leave you feeling?’) on a 5-point Likert scale (1, not at all pleased; 5, very pleased; see Supplementary Figure S2).

Assessment of pitch pattern processing. To provide a measure of elementary pitch pattern processing, we adopted a procedure similar to other tests of basic serial pitch discrimination originating in the classic work of Seashore (1919). We assessed participants’ ability to detect pitch direction changes in sequentially presented note pairs; the notes comprising each pair differed in pitch by either a tone or a semitone. Notes had piano timbre and duration 1 s with an inter-note gap of 1 s. Twenty trials (pairs) were presented and the direction of the pitch shift between notes was varied randomly across trials (10 ascending, 10 descending). The task on each trial was to decide if the second note was ‘higher’ or ‘lower’ than the first note.

Experimental procedure. Stimuli were presented from a notebook computer running Matlab via headphones (Audio-Technica) at a comfortable listening level (at least 70 dB) in a quiet room. Participants were first familiarised with each test using practice examples to ensure they understood the task and were able to comply reliably. Participant responses were recorded for offline analysis. During the tests no feedback was given about performance and no time limits were imposed. All participants completed the experimental tests comfortably in <30 min.

Analysis of behavioural data

All behavioural data were analysed using Stata12. Demographic characteristics and neuropsychological data were compared between participant groups using Fisher’s exact test for categorical variables, and for continuous variables, either two sample t-tests or Wilcoxon rank-sum tests where t-test assumptions were materially violated (e.g. due to skewed data distribution).

Logistic regression was used to analyse accuracy of melody and pitch direction decisions (correct vs incorrect) and pleasantness ratings (dichotomized based on a rating ≥3 (‘pleasing’) or <3 (‘not pleasing’) to avoid over-estimating the effect of gradations of pleasantness). For each analysis a global test was first used to jointly compare all groups, followed by pairwise group comparisons where a significant overall group effect was found. An interaction term was included to examine whether there was differential performance between groups for ‘finished’ vs ‘unfinished’ melodies. To take account of potentially confounding factors, age, gender, reverse digit span (an index of executive function and specifically, auditory working memory) and (in the
Table 1. General demographic, clinical and neuropsychological characteristics of participant groups

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Healthy controls</th>
<th>bvFTD</th>
<th>SD*</th>
<th>PNFA</th>
<th>AD</th>
</tr>
</thead>
<tbody>
<tr>
<td>General</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No. (m:f)</td>
<td>11: 11</td>
<td>9: 2</td>
<td>4: 2</td>
<td>2: 6</td>
<td>8: 6</td>
</tr>
<tr>
<td>Age (years)</td>
<td>68.5 (5.1)</td>
<td>65.8 (7.6)</td>
<td>66.2 (5.2)</td>
<td>71.5 (7.8)</td>
<td>68.6 (6.7)</td>
</tr>
<tr>
<td>Musical training (years)</td>
<td>4.5 (3.4)</td>
<td>4.8 (3.4)</td>
<td>4.3 (4.4)</td>
<td>3 (2.5)</td>
<td>4.4 (2.9)</td>
</tr>
<tr>
<td>Musical listening (h/wk)</td>
<td>9.7 (10.1)</td>
<td>5.5 (4.7)*</td>
<td>8.8 (8.6)*</td>
<td>5.5 (7.4)</td>
<td>8.6 (11.0)</td>
</tr>
<tr>
<td>Education (years)</td>
<td>16.7 (2.0)</td>
<td>15.3 (3.4)</td>
<td>14.2 (3.1)</td>
<td>16.9 (2.2)</td>
<td>15 (2.4)</td>
</tr>
<tr>
<td>MMSE (/30)</td>
<td>N/A</td>
<td>25 (3.8)</td>
<td>27 (2.5)</td>
<td>23 (9.5)</td>
<td>21 (5.0)</td>
</tr>
<tr>
<td>Symptom duration (years)</td>
<td>N/A</td>
<td>9.8 (5.5)</td>
<td>6.3 (1.8)</td>
<td>6.9 (3.7)</td>
<td>6.3 (1.9)</td>
</tr>
<tr>
<td>Neropsychological</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>General intellect: IQ</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>WASI verbal IQ</td>
<td>119 (7.0)</td>
<td>91 (16.5)</td>
<td>87 (11.5)</td>
<td>88 (16.1)</td>
<td>98 (14.4)</td>
</tr>
<tr>
<td>WASI performance IQ</td>
<td>121(10.6)</td>
<td>104 (15.5)</td>
<td>114 (19.1)</td>
<td>103 (18.9)</td>
<td>90 (21.5)</td>
</tr>
<tr>
<td>NART estimated premorbid IQ</td>
<td>122 (4.7)</td>
<td>108 (12.2)</td>
<td>107 (12.1)</td>
<td>104 (15.8)</td>
<td>113 (9.0)</td>
</tr>
<tr>
<td>Executive skills</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>WASI Block Design (/71)</td>
<td>46 (13.0)</td>
<td>33 (13.4)</td>
<td>41 (19.8)</td>
<td>21 (18.1)</td>
<td>18 (13.8)</td>
</tr>
<tr>
<td>WASI Matrices (/32)</td>
<td>25 (4.1)</td>
<td>21 (5.6)</td>
<td>24 (6.9)</td>
<td>20 (6.7)</td>
<td>13 (7.8)</td>
</tr>
<tr>
<td>WMS-R digit span forward (/12)</td>
<td>9 (2.1)</td>
<td>9 (2.5)</td>
<td>11 (1.5)</td>
<td>7 (2.0)</td>
<td>6 (2.2)</td>
</tr>
<tr>
<td>WMS-R digit span reverse (/12)</td>
<td>8 (2.0)</td>
<td>7 (2.7)</td>
<td>10 (2.2)</td>
<td>3 (2.2)</td>
<td>5 (1.7)</td>
</tr>
<tr>
<td>D-KEFS Stroop colour (s)</td>
<td>29 (4.2)</td>
<td>40 (10.3)</td>
<td>37 (10.2)</td>
<td>67 (20.9)</td>
<td>54 (22.5)</td>
</tr>
<tr>
<td>D-KEFS Stroop word (s)</td>
<td>21 (3.7)</td>
<td>27 (8.0)</td>
<td>22 (5.3)</td>
<td>52 (24.6)</td>
<td>37 (19.4)</td>
</tr>
<tr>
<td>D-KEFS Stroop interference (s)</td>
<td>58 (17.1)</td>
<td>81 (36.2)</td>
<td>62 (27.5)</td>
<td>149 (37.3)</td>
<td>107 (33.2)</td>
</tr>
<tr>
<td>Letter fluency (F: total)*</td>
<td>16 (5.5)</td>
<td>9 (4.6)</td>
<td>11 (4.9)</td>
<td>4 (2.7)</td>
<td>11 (5.0)</td>
</tr>
<tr>
<td>Category fluency (animals: total)</td>
<td>24 (5.4)</td>
<td>12 (3.8)</td>
<td>6 (2.8)</td>
<td>10 (3.4)</td>
<td>12 (5.4)</td>
</tr>
<tr>
<td>Trails A (s)#</td>
<td>32 (9.0)</td>
<td>45 (17.3)</td>
<td>35 (20.6)</td>
<td>69 (37.2)</td>
<td>73 (48.0)</td>
</tr>
<tr>
<td>Trails B (s)</td>
<td>73 (20.4)</td>
<td>148 (81.7)</td>
<td>89 (48.0)</td>
<td>233 (67.5)</td>
<td>175 (62.9)</td>
</tr>
<tr>
<td>WAIS-R Digit Symbol (total)</td>
<td>56 (10.7)</td>
<td>34 (8.7)</td>
<td>44 (11.2)</td>
<td>27 (12.0)</td>
<td>20 (15.6)</td>
</tr>
<tr>
<td>Episodic memory</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>RMT words (/50)</td>
<td>48 (1.8)</td>
<td>37 (8.8)</td>
<td>33 (6.6)</td>
<td>47 (3.7)</td>
<td>30 (5.8)</td>
</tr>
<tr>
<td>RMT faces (%)</td>
<td>44 (4.1)</td>
<td>33 (6.1)</td>
<td>32 (8.1)</td>
<td>37 (5.7)</td>
<td>33 (6.4)</td>
</tr>
<tr>
<td>Camden PAL (/24)</td>
<td>20 (2.5)</td>
<td>9 (6.6)</td>
<td>5 (4.5)</td>
<td>17 (4.5)</td>
<td>4 (4.0)</td>
</tr>
<tr>
<td>Language skills</td>
<td></td>
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<tr>
<td>WASI Vocabulary (/80)</td>
<td>71 (3.2)</td>
<td>50(18.7)</td>
<td>42 (17.6)</td>
<td>39 (17.9)</td>
<td>56 (10.0)</td>
</tr>
<tr>
<td>WASI Similarities (/48)</td>
<td>39 (4.8)</td>
<td>26(8.4)</td>
<td>22 (8.7)</td>
<td>28 (7.2)</td>
<td>26 (11.4)</td>
</tr>
<tr>
<td>GNT (/30)</td>
<td>26 (2.2)</td>
<td>8 (9.4)</td>
<td>0 (0.8)</td>
<td>17 (7.7)</td>
<td>16 (6.7)</td>
</tr>
<tr>
<td>NART (/50)</td>
<td>43 (3.6)</td>
<td>31 (11.5)</td>
<td>28 (10.5)</td>
<td>30 (12.8)</td>
<td>36 (7.2)</td>
</tr>
<tr>
<td>Single word repetition (/45)</td>
<td>N/A</td>
<td>N/A</td>
<td>45 (1.0)</td>
<td>33 (15.4)</td>
<td>N/A</td>
</tr>
<tr>
<td>Sentence repetition (/10)</td>
<td>N/A</td>
<td>N/A</td>
<td>10 (0.5)</td>
<td>6 (4.4)</td>
<td>N/A</td>
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<tr>
<td>Semantic memory</td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>BPVS (/150)</td>
<td>148 (1.9)</td>
<td>128(21.2)</td>
<td>120 (14.8)</td>
<td>144 (4.8)</td>
<td>145 (3.0)</td>
</tr>
<tr>
<td>Synonyms concrete (/25)</td>
<td>N/A</td>
<td>N/A</td>
<td>18 (2.6)</td>
<td>22 (2.9)</td>
<td>N/A</td>
</tr>
<tr>
<td>Synonyms abstract (/25)</td>
<td>N/A</td>
<td>N/A</td>
<td>17 (3.2)</td>
<td>21 (3.5)</td>
<td>N/A</td>
</tr>
<tr>
<td>Posterior cortical skills</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>GDA (/24)</td>
<td>16 (5.0)</td>
<td>13 (6.6)</td>
<td>12 (8.4)</td>
<td>5 (4.1)</td>
<td>5 (6.3)</td>
</tr>
<tr>
<td>VOSP Object Decision (/20)</td>
<td>19 (1.6)</td>
<td>17(1.9)</td>
<td>17 (2.7)</td>
<td>16 (5.6)</td>
<td>16 (3.4)</td>
</tr>
</tbody>
</table>

Notes: Mean (SD) scores are shown unless otherwise indicated; maximum scores are shown after tests (in parentheses). Bold denotes significantly different (P < 0.05) to the healthy control group.

AD, patient group with typical Alzheimer’s disease; BPVS, British Picture Vocabulary Scale (Dunn LM et al., 1982); bvFTD, patient group with behavioural variant frontotemporal dementia; D-KEFS, Delis Kaplan Executive System (Delis et al., 2001); GDA, Graded Difficulty Arithmetic (Jackson and Warrington, 1986); GNT, Graded Naming Test (McKenna and Warrington, 1983); MMSE, Mini-Mental State Examination score; N/A, not assessed; NART, National Adult Reading Test (Nelson, 1982); PAL, Paired Associate Learning test (Warrington, 1996); PIQ, performance IQ; PNFA, patient group with progressive non-fluent aphasia; RMT, Recognition Memory Test (Warrington, 1984); SD, patient group with semantic dementia; Synonyms, Concrete and Abstract Word Synonyms Test (Warrington et al., 1998); SD, semantic dementia; VIQ, verbal IQ; VOSP, Visual Object and Spatial Perception Battery (Warrington and James, 1991); WAIS-R, Wechsler Adult Intelligence Scale – Revised (Wechsler, 1981); WASI, Wechsler Abbreviated Scale of Intelligence (Wechsler, 1997); WMS-R, Wechsler Memory Scale - Revised (Wechsler, 1987).

Three patients (two with musicophilia) had predominantly left-sided temporal lobe atrophy, one (with musicophilia) had predominantly right-sided temporal lobe atrophy and two (also with musicophilia) had more symmetrical, bilateral temporal lobe atrophy.

Includes patients with musicophilia (five in SD group, one in bvFTD group).

*Time to complete Trails in seconds (maximum time achievable 2.5 min on task A, 5 min on task B) (Lezak et al., 2004).
Brain image acquisition and voxel-based morphometry analysis

Brain MRI data for voxel-based morphometry were acquired for 34 patients (12 AD, 11 bvFTD, 5 SD and 6 PNFA) on a Siemens Trio 3Tesla MRI scanner using a 32-channel phased array headcoil and a sagittal 3D magnetization prepared rapid gradient echo T1-weighted volumetric sequence (echo time/repetition time/inversion time = 2.9/2200/900 ms, dimensions 256 × 256 × 208, voxel size 1.1 × 1.1 × 1.1 mm). Volumetric brain images were assessed visually in all planes to ensure adequate coverage and to exclude artefacts or significant motion. Pre-processing of patient brain MR images was performed using the Segment routine and the DARTEL toolbox of SPM12 (Ashburner, 2007). Further details of imaging analysis are available in Supplementary Material. A study-specific mean brain image template, for displaying results, was created by warping all bias-corrected native space whole-brain images to the final DARTEL template in MNI space and calculating the average of the warped brain.

Using the framework of the general linear model, multiple regression was used to examine associations between regional grey matter volume and accuracy of melody classification as ‘finished’ or ‘unfinished’ (raw score), pleasantness rating of melodies (relative likelihood of rating unfinished melodies as ‘not pleasing’) and pitch direction task performance over the combined patient cohort. In separate design matrices, voxel intensity (an index of grey matter volume) was modelled as a function of each relevant musical behavioural characteristic, including syndromic group membership, age, gender, total intracranial volume and reverse digit span as nuisance covariates in all matrices. For each model, separate contrasts (one-tailed t-tests) assessed positive linear associations between grey matter and the parameter of interest across the combined patient cohort. Statistical parametric maps (SPMs) were thresholded at P < 0.05 after family-wise error (FWE) correction for multiple voxel-wise comparisons over the whole brain.

Results

General participant characteristics

Patient and healthy control groups did not differ in age (P = 0.41), gender (P = 0.16), educational background (χ² = 6.40, P = 0.20) or musical training (χ² = 1.51, P = 0.80) and syndromic groups had similar mean symptom duration (χ² = 2.59, P = 0.50). Patient groups showed the anticipated profiles of general neurological impairment (Table 1). Peripheral hearing function varied between participant groups [combined audiometric tone detection score, see Supplementary Material; overall F(4, 53) = 7.32, P < 0.001, Supplementary Table S1]); however, the absolute value of the difference was small and there was no significant correlation between peripheral hearing and accuracy on melody classification performance over the entire participant group (rho = −0.09, P = 0.50) nor within the combined patient cohort (rho = −0.08, P = 0.64).

Tonal expectancy processing

Group performance profiles on tonal expectancy tasks are summarized in Table 2 and in Supplementary Tables S2 and S3. Individual raw data are presented in Supplementary Figures S3–S5.

There was evidence of an overall group performance difference in the odds of correctly classifying melodies as ‘finished’ or ‘unfinished’ (P = 0.003; Table 2 and Supplementary Figure S3). Relative to the healthy control group, the bvFTD and AD groups each showed overall significantly less accurate classification of melodies (P < 0.05), driven principally by incorrect classification of ‘finished’ melodies (Table 2). The SD and PNFA groups did not show a significant deficit of melody classification. There were no significant differences between syndromic groups. Whereas healthy controls were equivalently accurate in classifying ‘finished’ and ‘unfinished’ melodies (odds ratio (OR) = 1.4 (95% confidence interval (CI) 0.7–2.9), P = 0.38), the AD group was significantly less accurate classifying ‘finished’ than ‘unfinished’ melodies (P < 0.001). Across the patient cohort, accuracy of melody classification did not correlate with prior musical training (rho = 0.19, P = 0.25) or auditory working memory (reverse digit span, rho = 0.20, P = 0.23) but was significantly correlated with a general measure of non-verbal executive function (WASI Matrices score, rho = 0.46, P = 0.003).

There was further evidence of an overall group difference in pleasantness ratings of ‘unfinished’ vs ‘finished’ melodies (P < 0.0001; Table 2 and Supplementary Figure S4). The healthy
control group was significantly more likely to rate ‘unfinished’ than ‘finished’ melodies as ‘not pleasing’ (OR = 7.7; CI 3.8–15.7); the ranges of individual raw healthy control ratings for ‘finished’ and ‘unfinished’ melodies overlapped (Supplementary Figure S4 and Supplementary Table S2), suggesting that controls did not simply rate melodies to align explicitly with their melody classification. Patient groups showed a qualitatively similar profile, in that all groups tended to rate ‘finished’ melodies as more pleasant than ‘unfinished’ melodies (Supplementary Figure S4). However, both the bvFTD and SD groups showed significantly less discrepant pleasantness rating profiles than did the healthy control group (P < 0.05) and the SD group was also significantly less likely than healthy controls to rate ‘unfinished’ melodies as ‘not pleasing’ (P < 0.001), while the PNFA and AD groups showed a profile comparable to healthy controls. Comparing disease groups revealed significant differences between the pleasantness rating profiles of the bvFTD and SD groups vs the PNFA and AD groups (P < 0.05). Patient group pleasantness rating profiles were similar for the complete melody set and when restricted to those melodies correctly classified as ‘finished’ or ‘unfinished’ (compare Supplementary Figures S4 and S5, see Supplementary Table S3): both the bvFTD and SD groups were significantly less likely than healthy controls to rate correctly classified melodies as ‘not pleasing’ (P < 0.05) while the PNFA and AD groups rated correctly classified melodies similarly to healthy controls.

**Pitch direction processing**

Relative to healthy controls, a deficit of pitch direction processing was evident in the bvFTD group (P = 0.03) and SD group (P = 0.03), but not the PNFA group (P = 0.33) or AD group (P = 0.48) (Table 2 and Supplementary Figure S3). Within the patient cohort, performance on the pitch direction and tonal expectancy tasks showed a borderline significant correlation (P = 0.05, rho = 0.31; analyses of tonal expectancy performance were adjusted for pitch direction score). Performance on the pitch direction task was significantly correlated with prior musical training (rho = 0.58, P = 0.0002) but not with executive measures (reverse digit span, rho = 0.26, P = 0.11; WASI Matrices score, rho = 0.28, P = 0.08).

**Neuroanatomical associations**

Significant neuroanatomical associations are summarized in Table 3 and SPMs are presented in Figure 2.

Significant associations between grey matter atrophy and impaired melody classification accuracy over the combined patient cohort were identified in right entorhinal cortex, anterior superior temporal gyrus and sulcus and medial orbitofrontal cortex, thresholded at P < 0.05FWE after correction for multiple voxel-wise comparisons over the whole brain. A significant association between grey matter atrophy and abnormal pleasantness rating of melodies (a tendency to rate unresolved melodies as less unpleasant than healthy controls) was identified in left inferior frontal gyrus (pars orbitalis), thresholded at P < 0.05FWE after correction for multiple voxel-wise comparisons over the whole brain. No significant grey matter associations of pitch direction processing were identified for the combined patient cohort at the prescribed threshold.

Discussion

Here we have shown that music models impairments of rule and reward processing in cardinal syndromes of FTD and AD. When compared with healthy controls, patients with bvFTD had both less accurate classification of melodies based on harmonic structure (impaired rule decoding) and altered affective responses to harmonic completion (abnormal reward valuation); patients with SD had abnormal reward valuation despite preserved rule decoding; patients with AD had normal reward valuation despite impaired rule decoding; and patients with PNFA showed a relatively normal profile. Taken together, these profiles argue for dissociable signatures of musical rule decoding and reward valuation across dementia syndromes.

Melody classification in the bvFTD and AD groups was impaired after taking elementary pitch pattern perception (performance on the pitch direction task) into account and classification accuracy did not correlate with auditory working
memory or prior musical training. It is therefore likely that the tonal expectancy test indexed a relatively specific impairment of musical rule processing. Such cognitive modularity may have contributed to the (on face value, somewhat surprising) preserved performance of the present PNFA group on melody classification, despite other evidence for impaired syntactical and pitch pattern deficits in this syndrome (Rohrer et al., 2012; Golden et al., 2017). On the other hand, tonal expectancy was aligned with general executive capacity, suggesting that this musical task may track disease severity. Abnormalities of musical reward valuation in bvFTD and SD were evident even in the context of correct rule decoding (for correctly classified melodies), further demonstrating that the processes of rule decoding and reward valuation are dissociable. The profiles of musical valuation exhibited by these syndromic groups were qualitatively similar to healthy controls despite quantitative differences, raising the possibility that more complex effects (such as biased ‘gain’ or altered precision of reward expectancy) might be revealed by larger cohort studies with scope to further refine the model of musical hedonic valuation in these diseases.

The musical profiles identified here resonate with behaviours exhibited by patients with FTD and AD in various other experimental and social contexts. Impaired rule-based processing on semantic categorization tasks has been demonstrated in bvFTD and AD (Grossman et al., 2001, 2003). More pertinently, these patients show impaired anticipation of future events based on previously experienced regularities (Irish et al., 2012; Irish and Piolino, 2016): a general mechanism for impaired decision-making. Patients with bvFTD show markedly impaired detection of rule violations in social scenarios (faux pas) and increased risk taking behaviour and reduced anticipation of future regret (aberrant reward prediction) (Rahman et al., 1999; Torralva et al., 2007; Bertoux et al., 2014; Bora et al., 2015). The present findings support previous evidence that detection of salience (rule violation) in socio-emotional contexts is generally impaired in bvFTD while perceptual and cognitive detection of salient events may be disengaged from affective reactivity in this syndrome (Sturm et al., 2006; Chiong et al., 2013; Clark et al., 2015b, 2016). In contrast, patients with AD have difficulty using rules to make decisions about future outcomes, but remain sensitive to affective outcomes (reward value) (Delazer et al., 2007; Dohnel et al., 2008; Sinz et al., 2008). AD and PNFA are substantially less likely than patients with bvFTD to exhibit strikingly aberrant responses to social rule violations (Bora et al., 2015; Clark et al., 2016), perhaps reflecting the relative extent to which affective awareness is preserved in these syndromes. In SD, decision-making based on modelling of future events depends critically on semantic function (Irish et al., 2011, 2012; Irish and Piolino, 2016): music (in contrast to most other rule-based systems) does not rely on extraneous semantic associations, perhaps accounting for the preserved ability to use musical rules exhibited by patients with SD here and in previous studies (Hallstone et al., 2009; Omar et al., 2010; Weinstein et al., 2011). Nevertheless (in line with the present profile), SD is frequently associated with abnormal valuation of biological and other rewards, including music (Fletcher et al., 2013, 2015a, b; Ahmed et al., 2014, 2015, 2016).

The neuroanatomical profiles identified here underline the involvement of brain networks mediating rule analysis, rule violation (salience) detection and reward evaluation in these.

Fig. 2. Statistical parametric maps (SPMs) of regional grey matter volume positively associated with tonal expectancy parameters in the combined patient cohort are presented (see also Table 3; t scores are coded on the colour bars. Grey matter associations of accuracy of melody classification signify musical rule decoding, indicated in blue; grey matter associations of melody pleasantness rating signify musical reward valuation, indicated in red-orange [see text for details]. SPMs are overlaid on coronal (left) and sagittal (middle, right) sections of the normalized study-specific T1-weighted mean brain MR image, selected to highlight right anterior superior temporal and entorhinal cortex (left), right medial orbitofrontal cortex (middle) and left inferior frontal gyrus pars orbitalis (right). SPMs are thresholded for display purposes at $P < 0.001$ uncorrected, however local maxima of areas shown were each significant at $P < 0.05$ after FWE correction for multiple voxel-wise comparisons over the whole brain (see Table 3).

Table 3. Summary of grey matter associations of tonal expectancy processing in patient cohort

<table>
<thead>
<tr>
<th>Region</th>
<th>Peak coordinate (mm)</th>
<th>Z score</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy of classifying melodies</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Entorhinal cortex</td>
<td>24 0 −50</td>
<td>5.22</td>
<td>0.008</td>
</tr>
<tr>
<td>Anterior superior temporal gyrus</td>
<td>48 3 −18</td>
<td>4.94</td>
<td>0.025</td>
</tr>
<tr>
<td>Medial orbitofrontal cortex</td>
<td>56 −10 −8</td>
<td>4.91</td>
<td>0.029</td>
</tr>
<tr>
<td>Anterior superior temporal sulcus</td>
<td>48 40 −22</td>
<td>4.92</td>
<td>0.028</td>
</tr>
<tr>
<td>Pleasantness rating of melodies</td>
<td>−51 33 −15</td>
<td>5.69</td>
<td>0.001</td>
</tr>
<tr>
<td>Inferior frontal gyrus (pars orbitalis)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes: The Table shows statistically significant positive associations between grey matter volume and accuracy of classifying melodies (‘finished’ vs ‘unfinished’) and pleasantness rating of melodies (likelihood of rating unfinished melodies as ‘not pleasing’), based on a voxel-based morphometric analysis of brain MR images for the combined patient cohort. Local maxima coordinates are presented with coordinates in MNI standard stereotactic space, thresholded at $P < 0.05$ after FWE correction for multiple voxel-wise comparisons over the whole brain. No significant associations were identified for the pitch direction task at the prescribed threshold.
diseases (Perry et al., 2014, 2015). Impaired tonal expectancy (classification of melodies) was associated with grey matter loss in right anterior superior and inferior temporal cortices and medial orbitofrontal cortex. In line with previous work (Peretz et al., 2001; Koelsch et al., 2006, 2013; Khalifa et al., 2008; Fujiwara and Cook, 2011; Hailstone et al., 2011; Omar et al., 2011; Hsieh et al., 2012; Salimpoor et al., 2013; Seger et al., 2013; Bonfiglio et al., 2015), this network may link cortical mechanisms mediating the structural analysis of melodies and harmonic hierarchies with paralimbic and orbitofrontal mechanisms mediating the cognitive representation and anticipation of musical emotion and reward. More generally, antero-medial temporal areas store previously learned knowledge and templates about sensory objects and regularities whereas prefrontal cortices implement and assess violations in rule-based algorithms (Michel et al., 2003; Christensen et al., 2011; Nazimek et al., 2013; Cohen, 2014; Gauvin et al., 2016; Henderson et al., 2016). Altered valuation of musical reward (affective rating of melodies) was associated with grey matter loss in left inferior frontal gyrus pars orbitalis. Functionally, this region behaves as a subdivision of lateral orbitofrontal cortex and links cortical mechanisms analysing hierarchical and rule-based patterns (such as linguistic and musical ‘syntax’) with mechanisms representing reward value (Belyk et al., 2017); it has been implicated previously in representing musical tension associated with violation of harmonic expectancy and anticipation of musical rhythms (Lehne et al., 2014; Vuust et al., 2014). We did not find neuroanatomical correlates in striatal or other subcortical structures previously implicated in processing musical reward (Omar et al., 2011; Salimpoor et al., 2013; Li et al., 2015). Previous studies have generally employed music familiar to the participants or explicit (e.g. monetary) valuation of music, perhaps implying that other motivational, emotional or subjective factors engage these subcortical mechanisms.

Impairments of social and emotional functioning remain difficult to assess in cognitively impaired patients, limiting the identification of relevant biomarkers and the design of effective interventions. Taken together, the present findings suggest that music may provide a useful paradigm for deconstructing complex behavioural disturbances in FTD and AD. Beyond its universality and relative tractability, music (as a dynamic stimulus that unfolds in time) inherently entails predictive coding: the anticipation of musical structure based on internalized models is a case of ‘aberrant rule and reward mechanisms in the dementias. We present findings provide a case for music as a useful probe of aberrant rule and reward mechanisms in the dementias. We propose a reappraisal of music as a unique window on complex behaviours in neurodegenerative disease.

Supplementary data
Supplementary data are available at SCAN online.

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Conflict of interest
None declared.

References


