Ventromedial frontal cortex mediates affective shifting in humans: evidence from a reversal learning paradigm

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Summary

How do the frontal lobes support behavioural flexibility? One key element is the ability to adjust responses when the reinforcement value of stimuli change. In monkeys, this ability—a form of affective shifting known as reversal learning—depends on orbitofrontal cortex. The present study examines the anatomical bases of reversal learning in humans. Subjects with lesions of the ventromedial prefrontal cortex were compared with a group with dorsolateral frontal lobe damage, as well as with normal controls on a simple reversal learning task. Neither form of frontal damage affected initial stimulus–reinforcement learning; ventromedial frontal damage selectively impaired reversal learning.

Keywords: frontal lobes; ventromedial prefrontal cortex; lesion; decision making; reward

Abbreviations: BDI = Beck Depression Inventory; CTL = age- and education-matched control subjects; DLF = dorsolateral frontal; fMRI = functional MRI; IADL = Instrumental Activities of Daily Living; MNI = Montreal Neurological Institute; VLF = ventrolateral prefrontal; VMF = ventromedial frontal

Introduction

An essential component of intelligent behaviour is the ability to learn the reward value of stimuli, which can change according to an animal’s circumstances and state. A previously rewarded stimulus may cease to be rewarding or even reverse its value and become punishing as a function of external changes in reinforcement contingencies or internal changes in appetite and satiety. Single cell recording and lesion studies have demonstrated that the orbitofrontal cortex is the substrate of flexible encoding of stimulus reward value in macaques. Orbitofrontal neurons encode the context-specific reward value of stimuli (Tremblay and Schultz, 1999; Rolls, 2000). Orbitofrontal lesions in monkeys impair ‘affective shifting’—the ability to adapt associative learning when an initially rewarded stimulus is no longer rewarded (extinction) or when the reward and punishment value of two stimuli switch (reversal learning). In contrast, lesions of dorsolateral prefrontal cortex do not affect either of these forms of learning (Jones and Mishkin, 1972; Dias et al., 1996).

Surprisingly little is known about the corresponding neural system in humans. Since the time of Phineas Gage, damage to ventral prefrontal cortex has been observed to result in personality changes, impaired impulse control and alterations in emotional and motivational state (Mesulam, 2002). It has been suggested that some of these changes are related to fundamental alterations in flexible stimulus–reward learning. Two recent lines of research have begun to investigate aberrant reinforcement processing in patients with ventral prefrontal damage, albeit from different perspectives. The first has focused on characterizing the impulsive decision-making that may follow VMF damage. It has been reported that patients with VMF lesions performing a complex gambling task are unable to change response strategies in the face of accumulating evidence that an initially preferred choice is detrimental in the long run (Bechara et al., 1997, 2000). Using simpler tasks that directly tested reversal learning and extinction, but in less well-characterized patients, Rolls and colleagues have argued more generally that the socially inappropriate behaviour of some frontally-damaged patients may reflect a basic inability to modify on-going behaviour in response to negative feedback (Rolls et al., 1994). Remarkably, however, functional imaging studies have yet to demonstrate a role for orbitofrontal cortex in the flexible stimulus–reinforcement
learning captured by reversal learning tasks; nor has reversal learning been assessed directly in humans with well-defined lesions of the ventromedial frontal (VMF) cortex.

An early study of this kind of stimulus–reward learning in brain-damaged humans—and the most informative to date—was carried out by Rolls and colleagues (Rolls et al., 1994). They examined the relationship between changes in day-to-day behaviour and reversal learning abilities in a heterogeneous group of patients. Subjects with primarily ventral frontal damage were impaired at both extinction and reversal learning, and had more severe disturbances of behaviour, when compared with those with damage outside this area. However, the anatomical specificity of this study was limited: many of the subjects did not have exclusive ventral frontal damage and the majority of the control group had non-frontal damage. This work therefore establishes the dissociability of reversal learning as a unique, frontally-mediated form of learning in humans. It does not, however, tell us what systems of prefrontal cortex participate in this process nor confirm the dorsal–ventral dissociation evident in animal work.

Two functional imaging studies have examined the neural basis of reversal learning in humans. An event-related functional MRI (fMRI) study of probabilistic reversal learning found activation in ventrolateral frontal (VLF) cortex when reversal error trials that preceded a correct response were contrasted with correct trials. Signal loss from the orbitofrontal cortex meant that its contribution to reversal learning could not be examined (Cools et al., 2002). A PET study with better ability to image orbitofrontal cortex observed no activation in this area when reversal learning was contrasted with other phases of a visual discrimination associative learning task (Rogers et al., 2000). More generally, however, several studies have implicated orbitofrontal cortex in reward processing, particularly in the context of uncertain or changing contingencies (Rogers et al., 1999; Elliott et al., 2000; O’Doherty, et al., 2000, 2001).

Technical limitations often make this region of the brain challenging to evaluate with standard fMRI techniques. Further, unlike lesion studies, imaging studies do not allow inferences to be drawn about whether a brain area has a necessary role in a given cognitive process. Therefore, we assess the neural substrates of flexible reinforcement processing in a study of human subjects with focal lesions of prefrontal cortex. Specifically, we test the hypothesis that injury restricted to the VMF lobes, but not to the dorsolateral frontal (DLF) lobes, impairs reversal learning in humans. We also examined the relationship between reversal learning performance and degree of functional impairment in subjects with frontal lobe damage in an effort to understand the influence of impaired affective shifting on day-to-day behaviour.

### Methods

#### Subjects

Subjects with fixed lesions involving the frontal lobes were identified through the research databases of the Hospital of the University of Pennsylvania and the Moss Rehab Research Institute. Any patient with a lesion confined to the frontal lobes, regardless of laterality, size or aetiology, was eligible. Patients were further classified as having a lesion that primarily involved either the VMF or DLF lobes (see below).

Exclusion criteria included past or intercurrent medical or neurological disease likely to impair cognition. Subjects were tested a minimum of 6 months after the brain injury. Age- and education-matched control subjects (CTL), free of neurological or psychiatric disease and not taking psychoactive medications, were recruited by advertisement in the local community. All participants provided written, informed consent in accordance with the Declaration of Helsinki; the study protocol was approved by the Institutional Review Boards of both participating centres.

Background information is provided in Table 1. There was no significant difference in age, level of education or premorbid IQ estimated by the National Adult Reading Test across the three groups of subjects. ANOVA (analysis of variance) demonstrated a significant effect of group on Beck Depression Inventory (BDI) scores \[ F(2,28) = 4.4, P = 0.02 \]. Post hoc Newman–Keuls tests showed that both patient groups had significantly higher scores than controls, but there was no significant difference between the two patient groups.

Four of eight subjects in the VMF group and five of 12 subjects with DLF damage were taking at least one psychoactive medication. These included specific serotonin uptake inhibitors, phenytoin, centrally-acting acetylcholinesterase inhibitors and methylphenidate.

Level of day-to-day function was assessed in two ways. Subjects with frontal damage (or, in three cases, their spouse or parent) completed a standard Instrumental Activities of Daily Living (IADL) scale (Gallo et al., 2000) to assess their day-to-day level of function. The scale was not completed for

### Table 1 Subject characteristics

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Age (years)</th>
<th>Sex (F/M)</th>
<th>Education (years)</th>
<th>NART IQ</th>
<th>BDI score</th>
<th>IADL score</th>
</tr>
</thead>
<tbody>
<tr>
<td>VMF</td>
<td>8</td>
<td>57 (12)</td>
<td>4/4</td>
<td>12.9 (2.0)</td>
<td>115 (10)</td>
<td>8.9 (8.1)</td>
<td>17.8 (3.4)</td>
</tr>
<tr>
<td>DLF</td>
<td>12</td>
<td>62 (10)</td>
<td>7/5</td>
<td>15.6 (2.7)</td>
<td>118 (11)</td>
<td>9.1 (3.3)</td>
<td>20.4 (1.4)</td>
</tr>
<tr>
<td>CTL</td>
<td>12</td>
<td>56 (16)</td>
<td>7/5</td>
<td>14.8 (2.7)</td>
<td>118 (12)</td>
<td>3.8 (2.6)</td>
<td>n/a</td>
</tr>
</tbody>
</table>

See text for details. Mean given with SD in parentheses. NART = National Adult Reading Test.
two subjects in the DLF group and one in the VMF group due to administrative problems. This scale evaluates seven domains, such as the ability to use the telephone and to manage finances. Total scores can range from 7 (dependent in all domains) to 21 (fully independent). Although this scale asks about very concrete day-to-day activities, patients with frontal damage may lack insight into their deficits. In order to corroborate the results of this functional self-assessment, a psychologist who knew the subjects well in her capacity as database manager (telephone contacts, arranging research visits, performing screening evaluations and, in several cases, visiting the subjects in their homes), but who was blind to the results of this study, independently classified subjects into one of three functional groups: (i) dependent on at least some IADLs; (ii) impaired, but not dependent; and (iii) independent, functioning at or very near pre-morbid level. The agreement in rank order between the two methods of measuring function was very good (Spearman $r = 0.84$, $P = 0.0003$). Given its greater range, the IADL score was adopted as the measure of function in further analyses. The VMF group was more impaired than the DLF group, as measured by the IADL score (Mann–Whitney $U$ test, $P = 0.02$).

Lesion analysis

Lesions were traced from the available CT or MRI onto the standard Montreal Neurological Institute (MNI) brain using MRICro software (Rorden and Brett, 2000). The subjects with frontal lobe injury were divided into two groups based on whether each lesion primarily involved the VMF or DLF lobes. Figure 1 shows the location and degree of overlap of the lesions in the two frontal groups. Lesions were due to rupture of anterior communicating aneurysms in seven out of eight VMF subjects. One subject had suffered an ischaemic stroke in the territories of both anterior cerebral arteries. DLF lesions were due to ischaemic stroke in eight cases, haemorrhagic stroke in three and tumour resection with local radiotherapy in one.

The lesions of the VMF group overlap maximally in the posteromedial orbitofrontal cortex and subjacent white matter. A region of interest that encompassed this area was outlined on the normalized brain on the basis of anatomical landmarks to permit more fine-grained correlational analysis. This region included the posterior half of the gyrus rectus, medial and middle orbital gyrus bilaterally, corresponding to the postero medial division of the orbitofrontal cortex according to the nomenclature of Hof (Hof et al., 1995).

A second analysis was performed to examine further the relationship between focal frontal damage and reversal learning performance, avoiding the constraint of a pre-defined region of interest. A lesion overlap image was constructed in which lesions were weighted according to the reversal learning performance of each subject, allowing direct visualization of the structure–function relationship.

Half the patients with dorsolateral involvement had lesions that extended to varying degree into the VLF cortex. Given the fMRI results implicating this region in reversal learning (Cools et al., 2002), the influence of such damage on both initial and reversal learning was also examined. A VLF region of interest was defined on the basis of these fMRI results using the following stereotaxic coordinates: $x = \pm 26$ to $\pm 50$; $y = +16$ to $+24$; and $z = -9$ to 8. The six DLF subjects with associated ventrolateral damage had a mean total lesion volume of 23.5 cc (SD 19), with a mean VLF involvement of 1.0 cc (SD 1.6). Five of the VMF subjects had very minor involvement of the VLF cortex [mean total lesion volume in this sub-group 14 cc (SD 10), mean ventrolateral involvement 0.1 cc (SD 0.1).
**Task**

Stimulus–reinforcement association learning and reversal learning were assessed by means of a simple computerized card game with play money stakes. Subjects were dealt two cards at a time from packs of different colours. One pack consistently concealed a $50 win, the other a $50 loss. Feedback was provided after each trial. After the learning criterion of eight consecutive cards chosen from the winning pack was met, the contingencies were switched. This constituted the reversal phase of the task. If the criterion was again met, the contingencies were switched again for a total of 50 trials, allowing up to five reversals.

**Statistical analysis**

The effect of group membership on errors was examined by repeated measures ANOVA, with learning phase (initial, reversal) the within subjects measure. Post hoc pairwise comparisons were made with the Newman–Keuls test. Correlations between continuous variables were assessed with Pearson’s r. Spearman rank-order correlations were determined for ordinal data. The significance level was set at $P < 0.05$.

**Results**

We tested stimulus–reinforcement associative learning and reversal learning in human subjects with damage either to the VMF lobes or DLF lobes, in addition to age- and education-matched control subjects. Errors were submitted to repeated measures ANOVA with group as the between subjects factor, and learning phase (initial, reversal) as the within subjects factor. This revealed significant main effects of group $[F(2,29) = 14.0, P < 0.0001]$ and of learning phase $[F(1,29) = 121.1, P < 0.0001]$, as well as a significant group × learning phase interaction $[F(2,29) = 7.3, P = 0.003]$.

As shown in Fig. 2, there was no significant difference across groups in the number of errors made while learning the initial discrimination $[F(2,29) = 0.61, P = 0.55]$. There was, however, a significant effect of group membership on the number of errors made when reversal learning was required $[F(2,29) = 20.2, P < 0.0001]$. Subjects with VMF damage made more than twice as many errors as normal controls in the reversal phase of the task. In contrast, the performance of subjects with DLF damage was indistinguishable from controls and significantly different from VMF performance (Newman–Keuls test, $P < 0.05$).

Of the measured demographic characteristics, only education was correlated with reversal learning performance across all subjects ($r = -0.42, P = 0.02$). The ANOVA examining the relationship between errors in the two learning conditions...
(initial, reversal) and group membership (CTL, DLF, VMF) was therefore repeated with level of education, dichotomized as high school or less (≤12 years) or any college (>12 years) included as a factor. The results were not substantially altered: effect of group \( F(2, 26) = 11.5, P = 0.0003 \); effect of learning condition \( F(1, 26) = 83.3, P < 0.0001 \); effect of group \( \times \) learning condition \( F(2, 26) = 6.2, P = 0.006 \); and effect of education \( F(1, 26) = 1.6, P = 0.22 \).

There was no significant correlation between the total volume of damaged tissue and reversal learning performance in the subjects with VMF damage \( r = 0.06, P = 0.81 \) nor between lesion volume and reversal learning performance across all subjects with frontal damage \( r = 0.08, P = 0.75 \). In contrast, the extent of damage to the posteromedial orbitofrontal cortex and its overlying white matter, a region that included the site of maximum lesion overlap, showed a significant positive correlation with degree of reversal learning impairment \( r = 0.72, P = 0.04 \) (Fig. 3).

In order to evaluate further the relationship between structural damage and reversal learning, a second lesion overlap image was generated for the seven of the eight VMF subjects with abnormal reversal learning performance. These seven subjects also had the worst performance of the frontal group as a whole \( (≥9 \text{ errors in the reversal phase}) \). They were divided into three groups on the basis of performance: 9 errors \( (n = 4) \); 12 errors \( (n = 1) \); or 15–16 errors \( (n = 2) \). Individual lesions were then given a weight of 1, 2 or 3 accordingly. The maximal overlap involved 10 of the 12 available layers, and occurred in the white matter in the posteromedial orbitofrontal cortex within a 16 mm depth from the orbitofrontal cortical surface (Fig. 4). Although the area of maximal overlap is on the left, there were too few subjects with unilateral right VMF damage to allow definite conclusions about laterality to be drawn.

The DLF subjects as a group had lesions scattered throughout the DLF lobes, with somewhat less overlap than the VMF group as a consequence. This raises the possibility that the normal reversal learning of the DLF group as a whole is masking an impaired subgroup with damage to a putative key DLF region. As mentioned, when reversal learning performance is examined on a case-by-case basis, the seven worst performers were VMF subjects. Only one DLF subject performed worse than any control, by a margin of a single error. This absence of outliers from the DLF group adds further support to the contention that damage to DLF cortex does not impair reversal learning.

Although this study was not originally designed to evaluate the role of VLF cortex in reversal learning, several subjects in the DLF group had lesions that extended into this region. We therefore performed a post hoc region of interest-based analysis to examine whether damage to the area of VLF cortex implicated in reversal learning studied with fMRI (Cools et al., 2002) influenced either initial or reversal learning performance. Table 2 shows the initial and reversal learning performance split by VLF involvement. Although data are presented for the VMF group split in this way, it should be noted that the degree of VLF involvement in the VMF group was minor (see Methods). The subgroups are small, but it does not appear that unilateral damage to VLF influences reversal learning. The only values for which the 95% confidence intervals do not overlap are the error rates for the DLF groups with and without accompanying VLF damage; the presence of VLF damage in the DLF group is associated with better initial learning performance. Three of the six subjects with damage that involved ventrolateral, but not ventromedial, prefrontal cortex had lesions of the right hemisphere; there was no indication of an effect of the laterality of VLF damage on either initial or reversal learning (data not shown).

Depression is a frequent sequela of damage to the frontal lobes (Robinson et al., 1984; Mayberg, 2001), and it has been argued that impaired decision making following frontal
Discussion

This study is the first to demonstrate that intact ventromedial prefrontal cortex, but not dorsolateral prefrontal cortex, is necessary for normal reversal learning in humans. This is not a general associative learning deficit: neither group of frontal patients was significantly worse than controls at learning the initial association in this simple task.

Although several of the subjects in the VMF group had damage extending into the medial frontal lobes in addition to orbitofrontal cortex involvement, correlational analysis indicates that the key area of damage leading to impaired reversal learning is the medial orbitofrontal cortex. More extensive damage to adjacent ventral and/or medial frontal structures, as quantified by the total lesion volume, did not correlate with worse performance. At least one selective lesion study in macaques has also shown that sustained impairments in reversal learning are specific to medial orbitofrontal cortex lesions (Iversen and Mishkin, 1970), although this finding has not been universal (Butter, 1969).

Similarly, additional damage extending into VLF cortex did not adversely affect either initial learning or reversal learning performance in either the VMF or DLF group. This finding contrasts with the fMRI study of Cools et al. (2002), in which right VLF cortex activation was found to be associated with reversal error trials. This VLF activity may be related to the additional demands of the more complex probabilistic reversal learning task used in that study. Alternatively, the putative role of this VLF region in reversal learning may be bilaterally represented, so that subjects with unilateral lesions remain unimpaired.

Orbitofrontal cortex dysfunction has been hypothesized to play a role in a variety of neurological and psychiatric conditions, including fronto-temporal dementia (Rahman et al., 1999), addiction (Volkow and Fowler, 1999), depression (Drevets et al., 2001) and obsessive-compulsive disorder (Nielen et al., 2002), to name a few. Behavioural methods to study the role of this frontal area in humans are limited; although VMF lobe damage impairs performance on a widely used gambling task (Bechara et al., 1997), recent work has called the anatomical specificity of this effect into question (Manes et al., 2002). Reversal learning tasks may provide useful tools for selectively probing orbitofrontal cortex function in such patient populations.

Lesion work in animals has demonstrated double dissociations between reversal learning (also termed affective shifting) and the ability to learn associations between reinforcement and new stimulus features for which the suppression of previous stimulus-reinforcement associations is not required (attentional shifting). Ventral prefrontal lesions impair affective shifts, but leave attentional shifts intact, while dorsal prefrontal lesions do the opposite (Dias et al., 1996). The concomitant experiment has yet to be reported in humans. Establishing the existence of such double dissociations would add considerably to the limited human literature in which lesion methods have been employed to study the separate processes mediated by dorsal and ventral prefrontal cortex (Bechara et al., 1998; Manes et al., 2002). The present work demonstrates a single dissociation for affective shifting, supporting the contention that this framework for understanding the anatomical basis of stimulus–reinforcement learning in animals is also of relevance in humans.

A previous study of reversal learning in humans with frontal lobe damage reported a correlation between reversal learning performance across all subjects with frontal damage and a global rating of the severity of behavioural change (Rolls et al., 1994). We also found that level of function—in

<table>
<thead>
<tr>
<th>Group</th>
<th>Learning errors mean (95% CI)</th>
<th>Reversal errors mean (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DLF not VLF (n = 6)</td>
<td>3.3 (1.2–5.4)</td>
<td>6.0 (4.5–7.5)</td>
</tr>
<tr>
<td>DLF and VLF (n = 6)</td>
<td>0.3 (0.0–0.87)</td>
<td>6.0 (4.4–7.6)</td>
</tr>
<tr>
<td>VMF not VLF (n = 3)</td>
<td>4.0 (1.1–6.9)</td>
<td>9.0 (9.0–9.0)</td>
</tr>
<tr>
<td>VMF and VLF (n = 5)</td>
<td>1.4 (0–6.1)</td>
<td>11.6 (6.4–16.8)</td>
</tr>
</tbody>
</table>

CI = confidence interval.
this case measured with a simple IADL scale—was negatively correlated with reversal learning impairment. Both these pieces of evidence raise the possibility that impaired affective shifting manifests itself in functionally important ways in everyday human behaviour, although experiments that identify the affected behaviours in more detail, and that move beyond correlational evidence, are required to explore this issue further.

The existence of neural circuitry specialized for rapidly unlearning or suppressing the influence of an established stimulus-reinforcement association is counter-intuitive. Our findings, adding to an extensive animal literature, argue that such a mechanism exists in humans and is, at least in part, anatomically distinct from the neural substrates that mediate the initial learning process.

Acknowledgements
We wish to thank Dr Marianna Stark for her help with subject recruitment and functional assessment. We also wish to thank the anonymous reviewers for their helpful commentary. This research is supported by NIH grants R01-AI41082, R01-DA14129, and NSF grant no. 0226060. L.K.F is supported by a Clinician-Scientist award from the Canadian Institutes of Health Research.

References

