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Selective Aspiration or Neurotoxic Lesions of Orbital Frontal Areas 11 and 13 Spared Monkeys’ Performance on the Object Discrimination Reversal Task

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Damage to the orbital frontal cortex (OFC) has long been associated with reversal learning deficits in several species. In monkeys, this impairment follows lesions that include several OFC subfields. However, the different connectional patterns of OFC subfields together with neuroimaging data in humans have suggested that specific OFC areas play distinctive roles in processing information necessary to guide behavior (Kringelbach and Rolls, 2004; Barbas, 2007; Price, 2007). More specifically, areas 11 and 13 contribute to a sensory network, whereas medial areas 10, 14, and 25 are heavily connected to a visceromotor network. To examine the contribution of areas 11 and 13 to reversal learning, we tested monkeys with selective damage to these two OFC areas on two versions of the ODR task using either one or five discrimination problems. We compared their performance with that of sham-operated controls and of animals with neurotoxic amygdala lesions, which served as operated controls. Neither damage to areas 11 and 13 nor damage to the amygdala affected performance on the ODR tasks. The results indicate that areas 11 and 13 do not critically contribute to reversal learning and that adjacent damage to OFC subfields (10, 12, 14, and 25) could account for the ODR deficits found in earlier lesion studies. This sparing of reversal learning will be discussed in relation to deficits found in the same animals on tasks that measure behavioral modulation when relative value of affective (positive and negative) stimuli was manipulated.

Introduction

Damage to the orbital frontal cortex (OFC) has long been associated with impairment in behavioral adaptation (Teitelbaum, 1964; Butter, 1969; Jones and Mishkin, 1972; Rolls et al., 1994; Dias et al., 1996; Bechara et al., 1997; Meunier et al., 1997; Ferry et al., 2000; Schoenbaum et al., 2002, 2003; Bohn et al., 2003; Chudasama and Robbins, 2003; Fellows and Farah, 2003; McAlonan and Brown, 2003; Hornak et al., 2004; Izquierdo et al., 2004; Kim and Ragozzino, 2005). This impairment is generally observed in the object discrimination reversal (ODR) task, in which subjects first respond to one of two cues to receive a reward and then inhibit responding to this rewarded cue upon reversal of the reward contingency. Following OFC damage, subjects typically learn the initial discrimination normally, but make more errors than controls on the reversals. For monkeys, the reversal deficit usually follows damage that encompasses several OFC subregions (Barbas and Pandya, 1989; Amaral et al., 1992; Carmichael and Price, 1994), including the middle areas 11/13, rostral area 10, ventromedial areas 14/25, and, in some instances, lateral areas 12/47 (Iversen and Mishkin, 1970; Jones and Mishkin, 1972; Meunier et al., 1997; Izquierdo et al., 2004). The one exception is the earlier study of Butter (1969), who demonstrated severe reversal deficits after lesions restricted to area 12, but not after lesions of either the anteromedial areas 10/11 or the posteromedial areas 13 and insular area (ia), suggesting that not all OFC subfields are critical for reversal learning [see also Roberts (2006) and Wallis (2007)]. These results are in line with recent anatomical maps and connectional networks of OFC subfields (Petrides and Pandya, 1984; Carmichael and Price, 1994; Price et al., 1996; Barbas et al., 2005) indicating the presence of two distinct networks: an “orbital” network (areas 11/13 and ia), and a “medial” network (areas 14/10) with its extensive connections with area 12 laterally and areas 25/32 medially (Barbas, 2007; Price, 2007). Functional distinctions between different OFC subfields came also from recent neuroimaging studies (Elliott et al., 2000a,b; Kringelbach and Rolls, 2003; O’Doherty et al., 2003; Hurliman et al., 2005). In the present study, we assessed the effects of selective lesions to OFC areas 11/13 on ODR in monkeys. Since reversal deficits following OFC lesions could be associated with task difficulty (Kim and Ragozzino, 2005), we also used two ODR tasks differing in the number of discrimination problems, i.e., one versus five problems, to be learned and reversed. Finally, as an operated comparison group, monkeys with neurotoxic amygdala lesions were also tested on the two tasks since these lesions do not disrupt ODR performance (Izquierdo and Murray, 2007). Given...
the earlier findings of Butter (1969), we predicted that restricting the OFC damage to areas 11/13 would spare reversal learning.

Materials and Methods

Subjects

Twenty adult male rhesus macaques (Macaca mulatta), ∼3.5 years of age at time of surgery, were used. All animals were housed individually, maintained on a 12 h light/dark cycle, and fed Purina primate chow supplemented with fresh fruit. Water was given ad libitum. During the behavioral testing, daily food intake was minimally restricted to ensure that the animal remained motivated to retrieve food rewards. The animals were divided into three experimental groups. Group C consisted of four sham-operated controls and two unoperated controls. Groups O-ibo and O-aso included two animals with neurotoxic lesions and five animals with aspiration lesions of areas 11 and 13, and group Aibo included seven animals with neurotoxic amygdala lesions. Before surgery, all animals received testing on social interactions, food preference, and emotional reactivity (intruder task). After surgery, they were given tests of novelty preference (visual-paired comparison task) and social interactions, and retrained on the food preference and intruder tasks. They were then tested on the devaluation reinforcer task, ∼12 months after surgery and received the one-pair and five-pair ODR, ∼18 months after surgery. The data obtained on social interactions, food preference, emotional reactivity (intruder task), and devaluation reinforcer task have previously been published (Machado and Bachevalier, 2006, 2007a and b).

Surgical procedures

All surgical procedures were performed under aseptic conditions, and anesthesia was induced by ketamine hydrochloride (10 mg/kg, i.m.) followed by isoflurane gas (1.0–2.0%, v/v, to effect). The animal’s head was secured into a nonferromagnetic stereotaxic apparatus (Crist Instrument), centered within the scanner bore (GE Signa 1.5 Tesla Echo Speed scanner, GE Medical Systems), and imaged with a 5" surface coil. Two MRI sessions (1–3 weeks before surgery and 7–10 d after surgery) were given to all animals except those in group C. During each session, two series of coronal images were taken through the entire brain: a T1-weighted structural (1 mm in thickness) and three Fluid Attenuated Inversion Recovery (FLAIR, 3 mm thick, each offset by 1 mm) scans. The presurgical T1-weighted images were used either to derive the stereotaxic coordinates for each injection site (Saunders et al., 1990) for animals in group A-ibo or to localize the orbital frontal sulci and determine the extent of orbital frontal areas 11 and 13 in animals of groups O-ibo and O-aso (Machado and Bachevalier, 2006, 2007a, b). Postsurgical FLAIR images were compared with matched presurgical FLAIR and T1-weighted images to accurately identify localized areas of edema indicative of neurotoxic-induced cell death, and were therefore used to quantify the extent of lesion for all animals in groups A-ibo and O-ibo (Máleková et al., 2001; Nemanic et al., 2002). Postsurgical T1-weighted images were compared with matched presurgical T1-weighted images to identify the location and quantify the extent of orbital frontal cortex aspiration lesions (group O-aso).

Magnetic resonance imaging

MRI procedures were performed under anesthesia induced by ketamine hydrochloride (10 mg/kg, i.m.) followed by isoflurane gas (1.0–2.0%, v/v, to effect). An intravenous drip of 0.45% sodium chloride was used for hydration, vital signs were continuously monitored, and a heating pad placed under the animal prevented hypothermia. Animals in group A-ibo were repositioned into the stereotaxic apparatus, whereas those in groups O-ibo and O-aso had their head secured into a head holder, which permitted free rotation of the animal’s head during surgery. The animal’s head was then shaved and disinfected with Nolvasan solution, and a local anesthetic (Marcaine, 25%, 1.5 ml, s.c.) was injected under the skin along the incision line. The skin and connective tissue were incised and gently retracted together with the temporalis muscles. Each group then underwent lesion-specific procedures.

Orbital frontal cortex lesion. Orbital frontal cortex lesions (both ibotenic and aspiration) were intended to damage the middle sector of the orbital frontal surface, including areas 11 and 13 (Barbas and Pandya, 1989; Amaral et al., 1992; Carmichael and Price, 1994). Given the individual variations in the shape and length of the orbital sulci, presurgical T1-weighted MR images were used to reconstruct the ventral surface of the frontal lobe for each animal. The boundaries of areas 11 and 13 on the
ventral surface of the frontal lobe were defined as (1) a line joining the anterior tips of the medial and lateral orbital sulci, anteriorly, (2) a line joining the medial bank of the lateral orbital sulcus to the olfactory stria just anterior to its division into the medial and lateral olfactory tracts, posteriorly, (3) the medial border of the olfactory stria, medially, and (4) the medial bank of the lateral orbital sulcus from its anterior tip to the posterior border of the lesion, laterally. These borders approximate the extent of areas 11 and 13 in the macaque monkey (Fig. 1, left panels).

The bone of the supraorbital ridge was opened and eroded, the dura was cut and retracted, and the brain was gently elevated to gain a full view of the orbital frontal surface. With the aid of a surgical microscope, the lateral and medial orbital sulci and the olfactory stria were visualized. For the neurotoxin injections, 17–36 injection sites were equally spaced (~2 mm apart in the lateral/medial and anterior/posterior planes) within these borders. A 30 gauge needle attached to a 10 μl Hamilton syringe by polyethylene tubing was used to manually inject 0.4–0.8 μl of ibotenic acid (0.4 μl/min) at each site. For the aspiration lesions (Mennier et al., 1997), 21 and 23 gauge aspirating probes in combination with electrocautery were used to inject 0.2–0.6 μl of ibotenic acid (Biosearch Technologies, 10 mg/ml in PBS, pH 7.4) into the appropriate injection coordinates. Two Hamilton syringes were filled with isotonic saline, replaced, and used to inject 0.2–0.6 μl of ibotenic acid to each site at a rate of 0.4 μl/min. To reduce intracranial swelling, Mannitol (20%, 1 ml/min, i.v.) was given just before the end of the surgical procedure.

Neurotoxic amygdala lesion. Using the presurgical T1-weighted MR images, the coordinates of 15 injection sites were selected within the amygdala to damage all amygdaloid nuclei. Two small craniotomies were performed to expose the brain just above the injection sites and small slits cut but no needle penetrations occurred. Each hemisphere was aspirated from the orbital frontal areas 11 and 13, as well as inadvertent tissue damaged found on all postsurgery T1-weighted images were drawn and used to inject 0.2–0.6 μl of ibotenic acid to each site at a rate of 0.4 μl/min. To reduce intracranial swelling, Mannitol (20%, 1 ml/min, i.v.) was given just before the end of the surgical procedure.

Sham lesions. For sham lesions, bilateral craniotomies (similar to those described above for the MR investigation.) were made as described above. The dura was cut but no needle penetrations occurred.

Following the specific surgical procedure of each type of lesion, tissues were closed in anatomical layers, and the animal was removed from isoflurane gas and recovered in the surgical facility until it could breathe on its own and maintained an SPO2 of >98% for 1 h. Beginning 12 h before surgery and continuing until 1 week after surgery, all animals were treated with dexamethasone sodium phosphate (0.4 mg/kg, i.m.) and Cephazolin (25 mg/kg, i.m.) to prevent excessive immunoreactivity and protect against infection, respectively. For 3 d following surgery, all animals received an agarose (aceticaminophen, 10 mg/kg, p.o.) to minimize pain. All animals were given a 7-d postsurgical regimen of dexamethasone sodium phosphate (0.4 mg/kg, i.m.) to reduce swelling, Cefazolin (25 mg/kg, i.m.) to minimize infection, and aceticaminophen (10 mg/kg, p.o.) for postoperative pain management.

Lesion assessment
Because eight animals (cases O-ibo-1 and -2, cases O-aso-1 and -2, and cases A-ibo-1 to -4) died in the flooding of Tropical Storm Allison in June 2001, the extent of lesion in those cases was determined using the presurgical and postsurgical scans (Malkova et al., 2001; Nemanic et al., 2002). Histology is available only for the six remaining animals (cases O-aso-4 to -6 and cases A-ibo-6 to -8) that were added to the study after the storm.

MRI lesion assessment. For all animals in groups A-ibo and O-ibo, presurgical T1-weighted 1 mm coronal images were matched with drawings of coronal sections from a normalized rhesus monkey template brain at 1 mm intervals. Hypersignals identified on FLAIR MR images were plotted onto corresponding drawings from the template brain, which were then imported into a Java-based image analysis program (Image; http://rsb.info.nih.gov/ij/) to measure the surface area (in pixels squared) of damage for intended targets, as well as all adjacent areas that may have sustained inadvertent damage. For any given region of interest (ROI), the surface area of hypersignals on each section through each hemisphere was summed and then multiplied by image thickness (1 mm) to calculate a total volume of damage (Gundersen and Jensen, 1987). For each ROI, the volume of damage for each hemisphere was then divided by the volume of that ROI in the normal brain to indicate a percentage of the total volume damaged.

For animals in group O-aso, presurgical and postsurgical T1-weighted 1 mm coronal images were matched to corresponding drawings from the normalized rhesus monkey template brain. The extent of orbital frontal tissue damaged found on all postsurgery T1-weighted images were drawn onto the corresponding drawings of the normal brain and extent of tissue aspirated from the orbital frontal areas 11 and 13, as well as inadvertent damage to adjacent cortical areas 10, 12, 14, 25, and ia were measured as described above for the MR investigation.

Histology lesion assessment. At completion of behavioral testing, the six animals were sedated (ketamine hydrochloride, 10 mg/kg, i.m.), given a lethal dose of sodium pentobarbital, and perfused intracardially with 0.9% saline followed by 4.0% paraformaldehyde. The brain was photographed and cut frozen at 50 μm in the coronal plane. Every fifth section was mounted and stained with thionin to visualize cellular bodies, and every 20th section was mounted and impregnated with silver (Gallyas, 1979) to visualize fibers. Histological sections were microscopically examined to evaluate the extent of cell loss, gliosis, and fiber damage. Percentage of damage to intended areas and inadvertent damage to adjacent areas was calculated as above by plotting the damage seen on each histological section onto a matched digital drawing of a coronal section from the template brain. For all cases that received both MR and histological investigation of the lesion extent, results obtained for each monkey with both procedures were then compared (Pearson’s moment correlation).

Behavioral testing
Apparatus and stimuli. Animals were trained in a Wisconsin General Testing Apparatus (WGTA) located in a darkened room containing a white noise generator to mask external sounds. The WGTA was equipped with a tray containing three food wells (2 cm in diameter, 1 cm deep, and 13 cm apart center to center). Only the two lateral wells were used to hide food rewards, i.e., M&M (Mars) or raisin (Sun-Maid Growers of California), under three-dimensional junk objects varying in color, shape, and texture.

Single-pair object discrimination reversal task. In this task (Jones and Mishkin, 1972), two objects formed a single discrimination problem.
Animals had first to learn which of the two objects was associated with the food reward (acquisition phase) followed by six reversals. During the first trial of the acquisition phase, both objects covered a food reward and the object selected by the animal became the rewarded object (S⁺) for the remaining trials of the phase. Left/right positions of the S⁺ object selected by the animal became the rewarded object (S⁺) trial of the acquisition phase, both objects covered a food reward and the S⁺ placed beside the empty well. This correction was repeated as many times as necessary until the animals displaced the S⁺. The number of times the correction trial was repeated provided a measure of perseverative errors (Jones and Mishkin, 1972; Meunier et al., 1997).

Five-pair object discrimination reversal task. Only a subset of animals in each group (cases C-4 to -7; A-ibo-7 to -9; O-asp-4 to -6) were given a more difficult version of the ODR task, consisting of five concurrent discrimination problems. Ten novel objects were selected to form five pairs with only one object serving as the S⁺ in each pair. Again, the S⁺ for each pair was selected on the first five trials when both objects of the pairs covered a reward. A total of 40 trials were given per day so that each pair was repeated eight times within a daily session in a pseudorandom order. Similar to the one-pair ODR, the five-pair ODR consisted of an acquisition phase followed by six reversals. Criterion was set at 37 correct choices in 40 trials (>90%) in one day followed by 34 correct choices in 40 trials (>85%) in the next day.

**Data analysis**

For both tasks, performance scores for the acquisition phase included (1) total number of trials given and (2) total number of errors made before reaching criterion performance. For each reversal and for six reversals combined, performance scores included (1) reversal errors as defined as errors committed in the first attempt of each trial and (2) perseverative errors, as defined as the number of times a trial was repeated before the animal selected the positive object during the correction procedure. Thus, scores for reversal errors and perseverative errors did not overlap.

First, for the one-pair ODR, we determined whether the type of orbital frontal lesions differently affected performance on the task, scores of group O-ibo were compared with those of group O-asp using Student’s t tests (SPSS v. 15). No differences between the two orbital frontal lesion techniques were detected for trials to criterion in the acquisition phase, reversal errors, and perseverative errors (one-tailed t = 0.576, df = 5, p = 0.59; t = 0.89, df = 5, p = 0.44; and t = 0.73, df = 5, p = 0.49, respectively). Therefore, for group comparisons, these two groups were combined into a single group designated group O in Results. Despite the similar effect of the two types of lesions, statistical analysis excluding the two cases with neurotoxic lesions that demonstrated sparing of some of the OFC fields 11 and 13 (see Tables 1 and 2 and Fig. 4) were also reported.

For both tasks, one-way ANOVAs were used for group comparisons on trials and errors in the acquisition phase and on the total number of reversal errors and perseverative errors obtained across the six reversals, using one-tailed tests. General linear model repeated-measures ANOVAs (SPSS v. 15) with group (3) and reversals (6) were used to compare performance across the six reversals. A Huynh-Feldt correction was used to adjust the degrees of freedom if sphericity could not be assumed. Significant main effects of group were investigated further using one-tailed tests. General linear model repeated-measures ANOVAs were corrected by rerunning the erroneous trial with the S⁺ covering the

### Table 3. Extent of intended and unintended damage in group A-ibo

<table>
<thead>
<tr>
<th>Cases</th>
<th>Amygdala</th>
<th>Hippocampal formation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L R Avg W</td>
<td>L R Avg W</td>
</tr>
<tr>
<td>A-ibo-1</td>
<td>20.6</td>
<td>82.2 51.4</td>
</tr>
<tr>
<td>A-ibo-2</td>
<td>48.9</td>
<td>88.1 68.5</td>
</tr>
<tr>
<td>A-ibo-3</td>
<td>27.1</td>
<td>73.1 50.1</td>
</tr>
<tr>
<td>A-ibo-4</td>
<td>79.1</td>
<td>92.5 85.8</td>
</tr>
<tr>
<td>A-ibo-7</td>
<td>80.8</td>
<td>96.4 88.6</td>
</tr>
<tr>
<td>A-ibo-8</td>
<td>29.6</td>
<td>44.3 36.9</td>
</tr>
<tr>
<td>A-ibo-9</td>
<td>43.9</td>
<td>75.2 59.6</td>
</tr>
<tr>
<td>X</td>
<td>47.1</td>
<td>78.8 62.9</td>
</tr>
</tbody>
</table>

Data are the estimated percentage of damage as assessed from MR (postsurgical T2 FLAIR) images. Abbreviations are as in Table 1.
post hoc Bonferroni tests and significant interactions between Group and Phase were investigated with one-tailed paired-sample t tests.

In addition to the aforementioned behavioral measures, we also conducted a general linear model repeated-measures ANOVA analysis of the distribution of errors according to the “stages of learning” previously proposed by Jones and Mishkin (1972). Briefly, these three stages are meant to assess potential group differences in learning patterns. Each stage is calculated by summing the cumulative correct responses per stage over the six reversals (not including the correction trials). Stage I is characterized by scores of 9 or fewer correct responses out of 30 trials, and represents the learning stage at which previous stimulus-reward associations must be suppressed. Stage II is characterized by scores between 10 and 20 correct responses out of 30 trials, and represents the period when the animal is performing at or around chance level. Finally, stage III is characterized by scores between 21 and 30 correct choices out of 30 trials, and represents the learning period in which the subject consistently associates the stimulus with the reward at a near criterion level.

To compare performance between the two tasks, multivariate repeated-measures ANOVAs (groups × reversals × tasks) were performed. We also conducted Pearson product moment correlation matrices (SPSS v. 15) to compare extent of lesion for groups A-ibo, O-ibo, and O-asp with reversal errors and perseverative errors. All unintended damage to surrounding areas > 5% per area was included in the analysis.

Results
Lesion extent
The extent of lesion based on MR images has been described in detail in previous reports (Machado and Bachevalier, 2006, 2007a,b, 2008). Tables 1–3 summarize the extent of intended and unintended damage for each animal of group O and group A-ibo, respectively. The weighted average (W) (Hodos and Bobko, 1984) was calculated to determine whether damage was highly unilateral (W < 25%) or particularly extensive and bilaterally symmetrical (W > 50%). Histological sections were available for three cases in each group (cases O-asp-4 to -6 and A-ibo-7 to -9) and comparisons between neuroimaging and histological estimation of lesion were performed for these six cases. These correlations were very high for both groups A-ibo and O-asp (Pearson, r = 0.935 and 0.910, respectively).

For group O, the aspiration lesions resulted in damage largely confined to areas 11 and 13, ranging from 83.3 to 95.9% as estimated from MRI (Tables 1, 2) and resulting in 78–94% cell loss from histological material in cases O-asp-4 to -6. Figures 1 (right panels) and 2 illustrate the extent of OFC lesion on the ventral view of the brain and on histological coronal sections for these three cases. Reconstructions of the OFC aspiration lesions on the ventral view of the brain and T1 MR images at three anterior–posterior levels are shown in Figure 3. In contrast, the two cases with neurotoxic OFC lesions received incomplete damage to areas 11 and 13 (range: 33.3–35.6%). Hypersignals were mostly confined to the superficial cortical layers and avoided the deepest layers, resulting in comparably lower total volume, but approximately the same extent of surface area damaged relative to group O-asp. Unintended damage for all cases (Tables 1, 2, Fig. 4) included area 10 (range: 0–12.3%), area 12 (range: 3.0–25.8%), area 14 (range: 1.0–36.4%), and ia (range: 8.3–32.9%).

For group A-ibo, the lesions resulted in damage largely confined to the amygdaloid nuclei, ranging from 36.9 to 88.6% as estimated from MRI (Table 3) and resulting in 27–73% cell loss from histological material in cases A-ibo-7 to -9. MR images (FLAIR) through the amygdala lesions are provided for two representative cases in Figure 5. In addition, histological sections through the amygdaloid lesions of case A-ibo-8, which received the smallest lesion of the group, are presented in Figure 6. Sparing in this case was limited to the ventrolateral portion of the amygdala. Unintended damage for all cases was mild and limited almost exclusively to the anterior hippocampus (average: 4%) (Table 3, Fig. 5, level +15).

One-pair ODR
The scores that each monkey obtained in the acquisition phase and six reversals are provided in Table 4 and illustrated in Figure 7.

Although two animals in group O and A-ibo required more trials than animals in group C to acquire the discrimination problem (Table 4), these differences did not reach significance ($F_{2,17} = 0.79, p > 0.05$ and $F_{(2,17)} = 0.98, p > 0.05$, for trials and errors, respectively).
Similarly, the three groups performed equally well across the six reversals as revealed by a nonsignificant group effect ($F_{(2,10)} = 2.72, p > 0.05$ and $F_{(2,10)} = 0.96, p > 0.05$, for reversal errors and perseverative errors, respectively). The reversal factor was significant for perseverative errors (reversal effect: $F_{(1,10)} = 4.22, p < 0.05$), but not for reversal errors (reversal effect: $F_{(1,10)} = 0.98, p > 0.05$). Post hoc analyses revealed that all animals made slightly, but significantly, more perseverative errors in reversal 1 compared with reversal 6 (one-tailed $t = 2.2, p = 0.040$). No other contrasts were significant, and the interaction between groups and reversals did not reach significance. The same findings were obtained when the two cases with neurotoxic OFC lesions were removed from the analyses (group effect: $F_{(2,15)} = 3.13, p > 0.05$ for reversal errors and $F_{(2,15)} = 1.56, p > 0.05$ for perseverative errors; reversal effect: $F_{(1,17,17)} = 5.94, p < 0.05$ for perseverative errors only). No other comparisons or interactions were significant.

Additionally, although number of total errors across the six reversals was slightly worse in groups O and A-ibo compared with group C (see Table 4), these differences did not reach significance for reversal errors or perseverative errors when cases with neurotoxic lesions were included ($F_{(2,17)} = 2.72, p > 0.05$ and $F_{(2,17)} = 0.96, p > 0.05$, respectively) or excluded ($F_{(2,13)} = 3.13, p > 0.05$ and $F_{(2,13)} = 1.56, p > 0.05$, respectively) from the analyses. It is interesting to note that two animals in group O (cases O-asp-5 and -6) made more perseverative errors than all other animals. These errors were made exclusively on the first two trials of the first reversal. Thus, case O-asp-5 made 89 perseverative errors on the first trial and few errors thereafter, and case O-asp-6 made 82, 175, and 11 perseverative errors in trials 1–3. However, a comparison of the number of perseverative errors made in the first day of reversal 1 showed again no group differences ($F_{(2,17)} = 0.54, p > 0.05$).

For stages of learning (Fig. 7), animals in the three groups did not differ ($F_{(2,17)} = 1.06, p > 0.05$), although the effect of stage was significant ($F_{(2,17)} = 36.25, p < 0.001$). Thus, all groups made more errors during stage II, compared with stage I and stage III ($F_{(1,17)} = 88.02, p < 0.05$ and $F_{(1,17)} = 4.27, p < 0.05$, respectively).

Finally, comparisons between scores on the task and percentage damage to adjacent cortical areas for group O, revealed a positive correlation between the amount of damage to area 14 and reversal errors made on the first reversal ($r = 0.78, p < 0.05$) but not with perseverative errors. This correlation was not significant when the two cases with neurotoxic OFC lesions were removed from the analyses (Tables 1, 2, Fig. 4).

### Five-pair ODR

As shown in Table 5 and Figure 7, the three groups learned the five-pair ODR at the same rate (group effect: $F_{(2,9)} = 0.40, p > 0.05$ and $F_{(2,9)} = 0.11, p > 0.05$, for trials and errors, respectively). Similarly, they performed equally well across the six reversals (group effect: $F_{(2,9)} = 0.75, p > 0.05$ and $F_{(2,9)} = 0.93, p > 0.05$; reversal effects: $F_{(2,9)} = 2.90, p > 0.05$ and $F_{(2,9)} = 1.64, p > 0.05$; group × reversal: $F_{(2,9)} = 0.84, p > 0.05$ and $F_{(2,9)} = 0.98, p > 0.05$, for reversal errors and perseverative errors, respectively). Overall performance across the six reversals was also comparable for the three groups ($F_{(2,9)} = 2.72, p > 0.05$ and $F_{(2,9)} = 0.96, p > 0.05$, for reversal errors and perseverative errors, respectively).

Furthermore, like for the one-pair ODR task, mean errors in the different stages of learning were similar for all groups ($F_{(2,10)} = 0.74, p > 0.05$), but differed between stages for all groups ($F_{(2,10)} = 8.67, p < 0.01$). Thus, all animals made more errors during stage II, than during stage I and stage III ($F_{(1,10)} = 14.51, p < 0.01$ and $F_{(1,10)} = 14.08, p < 0.01$, respectively) (Fig. 7).

Because only three animals in each group were tested on the five-pair ODR task, the sample size did not provide enough statistical power to determine whether or not a correlation existed between amount of damage and task performance.

### One-pair versus five-pair ODR performance

As shown in Figure 7, all animals obtained similar scores in learning the two tasks ($F_{(1,10)} = 0.16, p > 0.05$ and $F_{(1,10)} = 0.26, p > 0.05$ for trials and errors, respectively) and there were no group by task interactions. However, as predicted, all animals made more reversal errors in the five-pair ODR than the one-pair ODR.
(F(1,10) = 28.62, p < 0.001), but not more perseverative errors, and the interactions between group and task did not reach significance for any parameters. Similarly, for the stages of learning, only the effect of tasks was significant (F(2,10) = 8.27, p < 0.05; F(2,10) = 18.87, p < 0.05; and F(2,10) = 18.57, p < 0.05, for stages I, II, and III, respectively).

In summary, selective lesions of OFC areas 11 and 13 resulted in a remarkable sparing of abilities to solve the ODR task even when the animals were given a more challenging ODR task, in which they had to concurrently learn and reverse five discrimination problems. In addition, the lack of ODR impairment after selective lesions of the amygdala confirms and extends the earlier findings from Izquierdo and Murray (2004). Thus, neither OFC areas 11 and 13 nor the amygdala seem critical for suppression of responses that are no longer rewarded.

Discussion
Selective damage to the amygdala spares ODR task performance
Monkeys with selective amygdala lesions were unimpaired in the one-pair ODR task, confirming recent findings (Izquierdo and Murray, 2004) and extending these findings by showing that the lack of impairment persisted even when the task required the animals to concurrently learn and reverse five discrimination problems. Thus, the lack of impairment after amygdalectomy does not seem to result from the ease of learning a simpler reversal task. Nevertheless, amygdala neurons have been shown to modulate their activity based on shifts in reward contingency (Nishijo et al., 1988; Schultz, 2004; Paton et al., 2006; Salzman et al., 2007; Schoenbaum et al., 2007), suggesting an important role for the amygdala in computing and rapidly updating reward expectation. These two sets of findings do not necessarily conflict given that ODR performance is supported by a neural network that not only includes the amygdala but also the OFC areas and their connections with the medial temporal cortical areas and the striatum (Divac et al., 1967; Baxter and Murray, 2001; Clark et al., 2004; Budhani et al., 2007; Man et al., 2009). Thus, it is likely that other structures could allow nearly normal ODR performance in the absence of a functional amygdala. Future primate studies using reversible inactivation of the amygdala together with functional imaging could provide critical information on the interactions between the amygdala and other brain structures mediating reversal learning, as recently demonstrated in a neuroimaging study of two patients with amygdala damage (Hampton et al., 2007).

Selective damage to areas 11 and 13 spares ODR task performance
Damage restricted to OFC areas 11/13 alters neither the overall performance across the six reversals nor the learning patterns as measured by stages of learning. This lack of impairment persisted even when task difficulty was increased to five discrimination problems. However, it is still possible that more challenging task manipulation, such as the use of unpredictable reward outcomes providing greater ambiguity, may be needed to engage areas 11/13 as already shown in neuroimaging studies in humans (O’Doherty et al., 2001; Cools et al., 2002; O’Doherty et al., 2003).

This lack of impairment conflicts with previous reports in monkeys, but the differences cannot simply reflect procedural variations between studies. For example, a correction procedure was used in the present study to measure perseverative errors but not in the recent study of Izquierdo and Murray (2004). This correction procedure could have helped animals with OFC lesions to more rapidly learn that the reward contingency had changed. This possibility is unlikely, however, given that monkeys with OFC lesions in our earlier study (Meunier et al., 1997) were impaired on the ODR despite the use of a correction procedure. The major difference between this latter study and the present one is in the extent of the OFC lesion produced. In Meunier and colleagues (1997), the lesions included area 14 in addition to areas 11 and 13 as in the study by Izquierdo and Murray (2004). These data suggest either that the three OFC fields need to be damaged to yield the ODR deficit or that area 14 may play a more critical role in ODR learning than areas 11/13.
Second, Butter (1969) had demonstrated that ODR deficits did not occur after lesions restricted to either rostral OFC (e.g., 11 and 10) or caudal OFC (areas 13 and 1a), but were severe following selective area 12 lesions. Third, in humans, changes in activity related to response suppression necessary for ODR performance are found in area 12 and/or ventromedial areas (14, 25, and subgenual cortex) and less so in areas 11/13 (Bechara et al., 1999; Konishi et al., 1999; Elliott et al., 2000a; Fellows and Farah, 2003; Kringelbach and Rolls, 2003, 2004; Hurliman et al., 2005; Hampton et al., 2006, 2007; Budhani et al., 2007), and subgenual prefrontal regions are activated by negative feedback (Mars et al., 2005; van Duijvenvoorde et al., 2008; Zanoli et al., 2008). In addition, Kringelbach and Rolls (2004) have suggested that lateral OFC areas relate to punishers, whereas medial OFC areas relate to reward monitoring. Thus, given the anatomical connections between lateral area 12 and medial areas 14/25 (Barbas, 2007; Price, 2007), it is possible that, in the absence of areas 11/13, area 12 could still process the punishment associated with the object (no reward) and could pass this information directly to areas 14/25, which have been implicated in the visceral control of emotional regulation. Finally, the lack of ODR impairment after orbital frontal areas 11/13 strengthened other recent findings indicating that damage to the orbital frontal cortex might not affect all tasks requiring behavioral inhibition (Chudasama et al., 2007; Schoenbaum et al., 2007; Gaffan and Wilson, 2008) [but for an alternative explanation, see also Man et al. (2009)].

Yet, the significant changes in neural activity found in the same OFC areas 11/13 when reward contingency of stimuli has changed (Thorpe et al., 1983; Rolls, 1996; O’Doherty et al., 2001; Kringelbach and Rolls, 2003; Wallis and Miller, 2003) is at odds with the lack of ODR impairment following damage to these same areas. Although the loss of coding incentive value after orbital frontal lesions has been used to explain the deficits found in both the ODR and Reinforcer tasks, this proposal is no longer tenable in light of the present results.

**Amygdala and OFC interactions and the processing of affective values**

Despite normal performance in the ODR, the same animals with amygdala and areas 11/13 lesions were unable to shift choices away from objects associated with a devalued food in a reinforcer devaluation task (Machado and Bachevalier, 2007a). Given that earlier reports have stressed the critical contribution of amygdala/OFC interactions to flexibly adjust choice selection based on reward value as well as on reward contingency (Malkova et al., 1997; Baxter et al., 2000; Izquierdo and Murray, 2004, 2007; Izquierdo et al., 2004; Hampton et al., 2007; Schoenbaum et al., 2007), the present findings suggest that the interactions of the amygdala with areas 11/13 are necessary to adjust choices based on reward value (relative value) but less so for choices based on reward contingency (absolute value). This latter function could perhaps involve interactions between the amygdala and orbital frontal fields other than areas 11/13 since the amygdala is interconnected with several orbital frontal fields including the most medial subgenual fields and the most lateral area 12 (Barbas,

**Table 4. One-pair ODR task**

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<th>Group</th>
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Scores are total number of errors made before criterion days for the acquisition (Acq) and each of the six reversals (Rev 1 to Rev 6) as well as total reversal errors and total perseverative errors across the six reversals. X, Group mean.

This latter interpretation is consistent with several other findings. First, our correlation analysis already pointed to a positive association between the amount of damage to area 14 and perseverative errors. Second, Butter (1969) had demonstrated that ODR deficits did not
and total perseverative errors across the six reversals. In this regard, unilateral OFC fields and the amygdala may support performance on discrimination problems. The middle graphs depict the mean number of reversal errors for each stage of learning as determined by Jones and Mishkin (1972) (see details in Materials and Methods, Data analysis). C, animals with aspiration or ibotenic acid lesion of areas 11 and 13; O, groups O-asp and O-ibo, defined by Jones and Mishkin (1972) (see details in Materials and Methods, Data analysis). C, animals with aspiration or ibotenic acid lesion of areas 11 and 13; O, groups O-asp and O-ibo, defined by Jones and Mishkin (1972) (see details in Materials and Methods, Data analysis). A, animals with neurotoxic amygdala lesions; O-asp or O-ibo, animals with aspiration or ibotenic acid lesion of areas 11 and 13; O-asp-6 10 44 26 28 38 49 63 248 28 13 22 35 46 43 37 46 27 236 4 3 34 52 29 14 15 35 137 21 20 14 32 30 20 18 26 140 1 13 22 35 34 24 26 29 171 9

Table 5. Five-pair ODR task

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Scores are total number of errors made before criterion days for the acquisition (Acq) and each of the six reversals (Rev 1 to Rev 6) as well as total reversal errors and total perseverative errors across the six reversals. X, Group mean.

It is interesting to note that the reinforcer devaluation deficit after damage to the amygdala and areas 11/13 was also associated with difficulty in modulating emotional reactivity when the same animals were challenged with social stimuli differing in the magnitude of threat as assessed by the human intruder task (Machado and Bachevalier, 2007b). In this task, the animals are alone in a room into which an unfamiliar human entered, presenting either his/her profile to the animals (low intensity threat) or staring directly at the animals (high intensity threat). Before surgery, all animals showed higher frequency of tension-related behaviors in the staring condition than in the profile condition, indicating that they could modulate their emotional responses according to the intensity of threat provided by the intruder. After surgery, however, only sham-operated control animals continued to demonstrate this ability. Animals with amygdala and OFC lesions displayed higher, but similar, levels of tension-related behaviors to both conditions compared with their presurgical scores. Thus, as for the reinforcer devaluation task, both amygdala and ORB areas 11/13 seem critical to flexibly adjust emotional reactivity based on the magnitude of negative signals, such as threat.

The impairment in monitoring the positive and negative value of social cues after amygdala and OFC areas 11/13 lesions could also be the source of the striking behavioral changes found in the same animals when placed in small familiar social groups (Machado and Bachevalier, 2006). In this latter study, we reported that animals with amygdala lesions displayed several personality changes that preclude positive social interactions (increased exploration and excitability; decreased affiliation and popularity) and made abnormal responses to threatening social signals. In contrast, animals with orbital frontal lesions were involved in more aggressive interactions and responded abnormally to both affiliative and threatening signals. All together the data demonstrate that flexible decision-making mechanisms mediated by the amygdala and OFC areas 11/13 are critical to support normal social behavior. However, because these data were collected on animals that had been raised in large social groups but were singly housed at the time of these experiments, they will need to be confirmed by studies on animals maintained in a more naturalistic environment.

References
Baxter MG, Murray EA (2001) Impairments in visual discrimination learn-


