Functional dissociation between dorsal and ventral regions of the medial prefrontal cortex in rats

PASCALE GISQUET-VERRIER
Université Paris-Sud, Orsay, France

GORDON WINOCUR
University of Toronto, Toronto, Ontario, Canada

and

BENOÎT DELATOUR
Université Paris-Sud, Orsay, France

In order to investigate the respective roles of dorsal and ventral parts of the medial prefrontal cortex (mPFC) on a delayed alternation task, the acquisition of which is disrupted by large mPFC lesions (Winocur, 1991), rats with circumscribed lesions to the anterior cingulate (ACd) or the prelimbic-infralimbic (PL-IL) cortex were compared. As was predicted, ACd lesions severely impaired performance, and an analysis of the results suggests an involvement of this region in sequencing temporally ordered behavior. On the other hand, PL-IL lesions had no effect, a finding that was in contrast with previous evidence that PL-IL lesions induced delay-dependent deficits on a similar task (Delatour & Gisquet-Verrier, 1999). However, an important difference between the tasks related to the scheduling of critical delays. Whereas in the previous experiment, delay intervals were increased progressively in a block design, variable intertrial intervals were a constant feature of training in the present Experiment 1. In Experiment 2, rats with PL-IL lesions were administered Delatour and Gisquet-Verrier’s (1999) alternation task, but modified to incorporate Winocur’s (1991) variable-interval procedure that required less response adjustment. Under these conditions, the PL-IL group performed normally, as in Experiment 1. Taken together, the results provide evidence that the PL-IL cortex is implicated in processes that support attentional mechanisms and behavioral flexibility. Overall, the present results support the general hypothesis of subregionalization of mPFC functions.

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higher species (Preuss, 1995). On the other side, the ventral part of the mPFC, which is composed of the prelimbic (PL) and the infralimbic (IL) cortices (i.e., the PL-IL area), is involved in the limbic circuitry. The PL-IL area receives strong projections from the CA1/subiculum and entorhinal subfields of the hippocampal formation (Insauti, Herrero, & Witter, 1997; Jay & Witter, 1991), projects to the nucleus accumbens (Berendse, Galis-de Graaf, & Groenewegen, 1992), and has reciprocal connections with the amygdala (Ray & Price, 1992; Sesack et al., 1989).

In recent years, several investigators have examined the effects of selective lesions to the PL-IL and/or the ACd areas in a variety of behavioral paradigms. This work has given rise to various specific hypotheses regarding the functional significance of the respective regions. For example, Ragozzino, Adams, and Kesner (1998) have proposed that the ACd area is involved in working memory for egocentric space, whereas the PL-IL area is involved in working memory for allocentric space. More recently, the same authors have confirmed that these regions make different contributions to cognitive functions, with the PL-IL area, but not the ACd area, being involved in switching behavioral guiding strategies (Ragozzino, Wilcox, Raso, & Kesner, 1999). Earlier, Seamans, Floresco, and Phillips (1995) concluded that the PL cortex plays a critical role in prospective planning and that the ACd area is implicated in the flexible control of behavior that is necessary for selecting appropriate responses and altering strategies.

Work in our respective laboratories has also been concerned with this issue (e.g., Delatour & Gisquet-Verrier, 1996, 1999, 2000; Winocur, 1991, 1992; Winocur & Eskes, 1998). For example, Delatour and Gisquet-Verrier (1999) found that PL-IL lesions did not affect acquisition of a go/no-go conditional discrimination task but did induce a time-dependent deficit when delays were introduced between stimulus presentation and the opportunity to respond. By comparison, rats with large lesions to the mPFC that affected mainly dorsal regions were consistently impaired in conditional rule learning on similar tasks, even at very short delays (Winocur, 1991, 1992; Winocur & Eskes, 1998; see also Passingham et al., 1988). The combined data led Delatour and Gisquet-Verrier (1999) to suggest, in line with the views outlined above, that the dorsal mPFC contributes to the process of response selection, whereas the ventral mPFC participates in the organization of delayed responding.

A difficulty with work in this area is that conclusions often are based on studies that vary in numerous ways—most importantly, in terms of the location and size of lesions. Accordingly, one objective of the present study was to test the above hypotheses by comparing rats with circumscribed lesions that were restricted to the PL-IL or the ACd areas on a variable-interval (VI), delayed-alternation schedule (ITI). Successful performance on this task requires the rat to learn the basic response alternation rule and also be able to retain trial-specific information over short and long ITIs. We predicted a functional dissociation in which the ACd group would exhibit the deficit pattern observed by Winocur (1991; i.e., impairment in conditional rule learning). In contrast, in line with a working memory hypothesis, the deficit of the PL-IL group was expected to be more selective, affecting primarily performance at longer delays.

**EXPERIMENT 1**

**Method**

**Subjects**

Thirty-four male Long–Evans rats (50–57 days old), weighing 250–300 g at the start of the experiment, were obtained from the Trent University Breeding Center (Peterborough, Ontario). Rats were raised in social groups until they were 3–4 months old, at which time they were transported to individual wire-mesh cages and maintained on a 12:12-h day:night cycle with free access to food and water. Cages were located in a temperature-controlled room (21°C) in which the relative humidity was maintained at about 35%.

**Surgery**

The rats were assigned randomly to one of the three surgical groups: the PL-IL (n = 12), ACd (n = 11), and sham-operated (Sham, n = 11), groups. The rats were injected with atropine (i.p. injection, 0.3 ml) and were anesthetized 15 min later with pentobarbital (50 mg/kg/i.p.) before being mounted in a stereotaxic frame (Narashige Instruments, Tokyo, Model SR6). The skin on the skull was incised, and underlying tissue was removed. Horizontality of the head was verified, and holes were drilled around each injection's site. Ibotenic acid (Sigma Chemical Company, St. Louis) dissolved in phosphate buffered saline (10 mg/ml, final pH of 7.4) was injected bilaterally through a cannula (180 μm in diameter), using a microinjector. Bilateral lesions of the ACd cortex were made using the following injection coordinates (Paxinos & Watson, 1986): AP, +2.5 mm and +3.8 mm from the bregma; ML, ±0.5 mm from midline; and DV, −1.5 mm from the dura (each injection, 0.15 μl delivered over a 1-min period). Bilateral PL-IL cortical lesions were performed in a similar way at the following coordinates: AP, +3.6 mm; ML, ±0.7 mm; and DV, −3.4 mm (each injection, 0.5 μl delivered over a 4-min period). After each injection, the cannula was left in place for 5 min (ACd rats) or 10 min (PL-IL rats). Sham-operated rats were subjected to the same surgical procedure, but without penetration of the cannula. At the end of the surgery, the incision was sutured, and sulfamides were locally applied.

**Histology**

At the conclusion of behavioral testing, the rats were overdosed with pentobarbital (120 mg/kg) and perfused transcardially with saline, followed by a 10% buffered formalin solution. Brains were postfixed for one night in buffered formalin. After cryoprotection by immersion in a 30% sucrose–formalin solution for 48 h, the brains were cut on a freezing microtome into 40-μm sections. Every second section was taken, mounted on a gelatin-coated slide, and stained with cresyl violet.

Evaluation of the lesion size was performed by digitizing the lesion extent with a graphic software (Canvas, Deneba). To be included in the lesioned groups, it was necessary that cortical lesions, in each hemisphere, be damaged in more than 40% of the pregen-
tial lesion of the pregenual part of the ACd cortex. Only restricted encroachments into the dorsal PL cortex were noted and, on the dorsal extent, the medial precentral cortex was generally spared. Five rats were rejected owing to extensive invasion of the PL cortex. The remaining 6 rats sustained a restricted and subtotal ACd lesion (mean damage, 61.61% ± 6.28%; minimum, 40.36%; maximum, 85.41%).

Prelimbic-infralimbic cortical lesions. The rats in this group generally presented a bilateral lesion of most of the PL-IL cortices. One rat sustained damage to the medial orbital cortex and was retained. The dorsomedial PFC cortex (ACd and medial precentral cortices) was almost always preserved in all the rats. Five rats were discarded from behavioral analyses owing to very small lesions (n = 2), unilateral lesions (n = 2), or lesions extending too far caudally and ventrally in subcallosal regions (n = 1). The 7 remaining rats presented bilateral lesions centered on the PL-IL area (mean damage, 66.27% ± 4.24%; minimum, 59.76%; maximum, 82.32%).

Behavior
Acquisition of the variable-interval delayed alternation task. The results obtained during training are illustrated in Figure 2 in the form of no-go–go (Δ) latency ratios. Ratios were calculated and then averaged for each of the six ITIs. A high ratio would result from shorter latencies in the go trials than in the no-go trials. Thus, improved performance on the task over test sessions corresponded to progressive increases of the Δ ratio.

An analysis of variance (ANOVA), with one between-group variable (lesion) and one within-group measure (block), was performed on the Δ latency ratios obtained for the three experimental groups over the seven blocks of three training sessions. This analysis yielded a significant main effect of lesion [F(2,21) = 6.10, p < .01], as well as a significant main effect of block [F(6,126) = 14.11, p < .001], with no lesion × block interaction [F(12,126) = 1.66, n.s.]. Complementary analyses revealed progressively improved performance for each of the three experimental groups [Sham, F(6,60) = 5.45, p < .001; PL-IL, F(6,36) = 8.23, p < .001; ACd, F(6,30) = 4.22, p < .005]. The ACd-lesioned group, however, exhibited significantly lower Δ latency ratios than did the sham-operated group [F(1,15) = 18.30, p < .001] and the PL-IL-lesioned group [F(1,11) = 4.53, p = .053], whereas the PL-IL and Sham groups did not differ from each other (F < 1). Paired comparisons indicated that the ACd group had significantly lower Δ latency ratios than the Sham group on Blocks 2, 3, 5, and 7 (0.05 > ps < .01) and the PL-IL group on Blocks 2, 3, and 7 (0.05 > ps < .01).

Analysis of the last training block. The data for the final block were analyzed separately to assess various effects after performance on the task had stabilized. The go and the no-go latencies at each ITI for the seventh block of three test sessions are illustrated in Figure 3. A three-way ANOVA (lesion × type of trial × ITI) performed on response latencies indicated a significant main effect of
Figure 1. Reconstruction of PL–IL (left part) and ACd (right part) cortical lesions at different rostrocaudal levels (+4.7 to +1.7 mm from bregma) from the rats trained in Experiment 1. Black area, region lesioned for 75%–100% of the rats; hatched area, region lesioned for 50%–75% of the rats; dotted area, region lesioned for 25%–50% of the rats. PL–IL lesions were centered around the ventral medial prefrontal cortex, with minor encroachments on other prefrontal cortical areas. ACd lesions were restricted to this area and did not invade adjacent PL cortex. Nomenclature is adapted from Krettek and Price (1997). PrCm, medial precentral cortex; ACd, dorsal anterior cingulate cortex; ACv, ventral anterior cingulate cortex; PL, prelimbic cortex; MO/VO, medial orbital and ventral orbital cortices; VLO, ventrolateral orbital cortex; LO, lateral orbital cortex; IL, infralimbic cortex; dp, dorsal peduncular cortex; tt, taenia tecta; Accb, accumbens nucleus; Cpu, caudate putamen. From The Rat Brain in Stereotaxic Coordinates (plate 5 to plate 11), by G. Paxinos and C. Watson, 1986, San Diego: Academic Press. Copyright 1986 by Academic Press. Adapted with permission.
Figure 2. Delayed go/no-go alternation task in Experiment 1. Latency ratios (no-go latencies minus go latencies in seconds; mean ± SEM) of sham-operated rats (Sham), prelimbic–infralimbic-lesioned rats (PL–IL), and dorsal-anterior-cingulate-lesioned rats (ACd). Ratios were calculated for seven blocks of three consecutive sessions, independently of the delays. ACd rats showed a performance impairment throughout training relative to both Sham and PL–IL rats. Between group comparisons: ACd versus Sham, \(*p < .05, **p < .01, ***p < .001; ACd versus PL–IL, \(\times p < .05, \# \times p < .01\).

Discussion

The results of Experiment 1 indicated that ACd lesions impaired the acquisition of a go/no-go alternation task when trained under a variable ITI procedure. This disruptive effect was not modulated by the length of the ITI and seems to result from difficulties in withholding lever-pressing during the no-go trials. Moreover, the effect was not due simply to a slight delay in learning the task, since it was still present at the end of testing (i.e., after 21 sessions). The behavior of those rats with ACd lesions, which was very similar to that of the rats with large dorsal mPFC lesions in Winocur’s (1991) study that involved the same task, underscores the importance of this region for learning and performing the go/no-go alternation task.

In contrast, PL–IL lesions did not affect performance of the delayed alternation task at any delay. This result was surprising in that such lesions consistently produced delay-dependent deficits on similar behavioral tasks, including a conditional discrimination test involving go/no-go responding (Delatour & Gisquet-Verrier, 1996, 1999, 2000). This finding appears to challenge the hypothesis that this region is involved in working memory.

There are at least two possible explanations for this unexpected outcome. First, there may have been some idiosyncratic elements in the present task that allowed the PL-IL rats to respond normally even at relatively long delays. It has been suggested that, in some circumstances, rats can adopt alternative strategies that may prevent a delay-dependent deficit to occur (see Chudasama & Muir, 1997). Alternatively, the result may be due to specific training and design-related features of the present experiment. In their previous studies, Delatour and Gisquet-Verrier (1996, 1999, 2000) used a block design in which rats were trained on a given task with minimal
able delays throughout testing, this was an invariant part of the procedure that required no special adjustment.

For the following experiment, it was decided to test the effects of PL–IL lesions on a variation of the go/no-go conditional discrimination task that revealed clear delay-dependent impairment when a block design was used (Delatour & Gisquet-Verrier, 1999). The aim of Experiment 2 was to investigate whether such a delay-dependent deficit can still be evidenced when a variable-delay schedule, similar to that followed in Experiment 1, was adopted. In this test, conducted in operant chambers, rats had to learn to leverpress in response to a light stimulus in order to obtain food reward and to refrain from pressing in the presence of a tone in order to avoid shock. A variable delay (0–20 sec) between stimulus presentation and the appearance of a retractable lever was introduced in the first session and maintained throughout testing. If the deficit observed by Delatour and Gisquet-Verrier (1999) in their progressive delay paradigm was due to an inability to adjust to increasingly demanding changes to the response requirements, it is reasonable to expect that rats with PL–IL lesions would have less difficulty with the present task, which makes fewer demands on such processes. Such an outcome would be consistent with Delatour and Gisquet-Verrier’s (1999) suggestion that a working memory hypothesis is insufficient to account for PL–IL functions.

**EXPERIMENT 2**

**Method**

**Subjects**

Twenty-one male Sprague-Dawley rats were assigned to one of two groups: sham-operated rats (Sham, \( n = 9 \)) and PL–IL-lesioned (PL–IL, \( n = 12 \)) groups, using the same surgical and histological procedures as those described for Experiment 1. Housing conditions were also identical to those in Experiment 1.

**Apparatus**

Two operant chambers (Campden Instruments, U.K., Model 410) enclosed in sound-attenuated boxes (Model 412) were used for this experiment. Chambers were constructed of aluminum, with the exception of a Plexiglas aperture panel. The grid floor was made of 16 stainless barrels (0.5 cm in diameter) spaced 1 cm apart and was connected to a shock generator that could deliver a scrambled footshock. The left panel of each chamber was equipped with a central food magazine coupled to a 45-mg Noyes food pellets dispenser (Model 441) and a retractable lever, 10 cm to the right of the food tray. A 2-W bulb was placed 10 cm above the food magazine, and a speaker, through which a tone (1000 Hz, 80 Db) could be emitted, was mounted on the ceiling of the cage. Lighting of the operant chambers was provided by a 2-W houselight mounted on the ceiling, and a masking noise (about 40 dB) was provided by a ventilation fan placed on the right wall of the sound-attenuated boxes. Programming of experimental contingencies and data acquisition were accomplished by means of a PC. For each animal, data were printed and saved on magnetic disks.

**Procedure**

During pretraining and training, the operant chambers were continuously illuminated with the houselight.

**Pretraining.** The rats in each group were equally distributed among the two cages. On Day 1, the rats were placed in the chamber (with the lever retracted) for a 30-min period, during which two
food pellets were automatically delivered, every 60 sec, in the food magazine. On Day 2, the rats were trained on a CRF schedule. They were placed in the chamber with the lever inserted and coated with a gruel of crushed wet food pellets. During CRF, each leverpress was reinforced by two food pellets. The session ended after 50 leverpresses or after a 30-min period, whichever came first. Those rats that failed to press 50 times in the limited period of 30 min were given a second CRF session at the end of the day. On Days 3–4, leverpresses were reinforced on a 20-sec VI schedule during a 30-min daily session.

Training. Training in the VI delayed conditional discrimination task began on Day 3. At the beginning of each session, the rat was placed in the chamber with the lever retracted. After a 20-sec period, a 5-sec tone or a 5-sec light stimulus was delivered. Immediately following removal of the stimulus, a VI (0, 5, 10, or 20 sec) occurred, followed by the insertion of the lever in the chamber. A leverpress, within a 10-sec period, triggered the retraction of the lever and produced the reinforcer associated with that particular conditional stimulus. Following the light stimulus, a leverpress produced two food pellets, whereas a leverpress following the tone stimulus led to a footshock delivered through the grid floor. The next trial was then initiated (20 - n) sec later, where n (ranging from 0 to 10 sec) corresponded to the response latency to press the lever during the previous trial. When no leverpress was emitted within a 10-sec period, the lever was retracted, and the next trial was initiated 10 sec later.

Each daily session included 56 trials for a total duration of 28 min. Mild footshocks (0.15 mA) were administered during the first training session to punish responses to the tone stimulus. In subsequent sessions, the shock intensity was progressively increased until the rats consistently avoided pressing the lever in response to the tone stimulus. The rats were thus trained with different shock intensities ranging from 0.15 mA up to a maximal level of 0.5 mA. The light/tone sequences were pseudorandomly determined according to one of four preplanned sequences. In every session, there were 28 light and 28 tone trials, each stimulus being delivered a maximum of three consecutive times. For each session, each of the four different delays was randomly presented seven times following tone stimuli and seven times following light stimuli.

The number of responses to the light and tone stimuli in each session was recorded. To perform successfully, the rats had to learn to press the lever following the light (go-trials) and to avoid pressing the lever following the tone (no-go trials). Two types of error were defined and analyzed at the end of each session: (1) inhibition type error (pressing the lever following the tone, which led to a footshock) and (2) omission type error (omitting to press the lever following the light, which resulted in failure to obtain food reinforcement).

The rats were trained until they reached a criterion of performance for each delay condition. The criterion was defined as two consecutive training sessions in which the percentage of responses to the light was 50% higher than the percentage of responses to the tone, providing that the rats responded to the light on more than half of the light trials. The rats received at least 20 test sessions. Those that did not reach the criterion at all four delay intervals within the first 20 sessions received supplementary training until that criterion was achieved.

Results

Histology

PL–IL cortical lesions produced in the rats in Experiment 2 were very similar in terms of location and extent to those described in the first experiment. Seven rats were discarded because their lesions were considered too small. For the 5 remaining rats, lesions of the PL and the IL were extensive (global mean damage, 63.10% ± 8.96%; minimum, 57.78%; maximum, 87.4%) but largely spared the dorsomedial PFC. The orbitofrontal cortex was marginally damaged in 3 rats, but the encroachment was very small and not considered sufficient to justify exclusion.

Behavior

Acquisition of the variable-interval delayed conditional task. The responses made during the positively reinforced light trials (go-trials) and the negatively reinforced tone trials (no-go trials) are reported, in percentage terms, in Figure 4. Progressive differences in responding were observed between the go and the no-go trials through the successive four blocks of five training sessions. Differential responding for each type of trial was more prominent for the shortest delays than for the longest delays that separated the conditioned stimuli from the responses. An ANOVA performed on responses to tone and light stimuli with one between-group measure (lesion) and three within-group measures (type of stimulus, delay, and block) did not reveal any effect of lesion [F(1,12) = 1.47, n.s.] but did reveal a significant main effect of block [F(3,36) = 4.37, p < .05], type of stimulus [F(1,12) = 71.13, p < .001], and delay [F(3,36) = 15.33, p < .001].

Subsequent analysis indicated that, over the test sessions, the rats in both groups progressively increased their responses to the light stimulus [F(3,36) = 4.93, p < .01] and concurrently reduced their responses to the tone stimulus [F(3,36) = 28.45, p < .001]. This was confirmed by a significant block × stimulus interaction [F(3,36) = 80.80, p < .001], which was not affected by the lesion factor (F < 1).

Responses to both tone and light stimuli were affected by delay [F(3,36) > 7.84, ps < .001], and a significant delay × stimulus interaction [F(3,36) = 58.57, p < .001] indicated that the delays differentially affected responses to each stimulus. More precisely, a delay effect was observed for tone trials for all of the four training blocks [F(3,36) > 3.05, ps < .05], whereas performance on light trials was affected by delay only during Blocks 2 and 3 [F(3,36) > 3.21, ps < .05], suggesting a stronger effect of the delays on responses to tone than on responses to light.

Further analyses showed a significant lesion × delay interaction [F(3,36) = 3.25, p < .05], suggesting that the two groups differed in their sensitivity to the delays. This interaction resulted from the PL–IL rats’ exhibiting a significant delay-dependent effect between the second and fourth testing blocks [Fs(3,12) > 3.5, ps < .05], whereas Sham rats demonstrated a delay-dependent effect only during the last two conditioning blocks [Fs(3,24) > 4.66, p < .05].

Overall, PL–IL rats were able to perform the task adequately at all the delays. Between-group analyses performed at each ITI did not reveal any effect of lesion [Fs(1,12) < 2.87, n.s.]. Furthermore, the lesion × stimulus and the lesion × block interactions were not signifi-
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Figure 4. Delayed go/no-go conditional task in Experiment 2. Mean percentages of responses (±SEM) of sham-operated rats (Sham, n = 9) and prelimbic-infralimbic-lesioned rats (PL-IL, n = 5), during the go trials (light → food pellet: Sham, —○--; PL-IL, —■—■) and the no-go trials (Tone → electrical footshock: Sham, —●--; PL-IL, —■—■). Differences between the percentages of responses to the light and to the tone increased during the four blocks of five training sessions in every delay condition (0, 5, 10, and 20 sec). Response accuracy diminished when delays between the conditional stimulus and the response increased, with no difference between the Sham and the PL-IL groups.

...cant (Fs < 1), indicating that the lesion did not affect performance on the conditional task, regardless of the length of the delay.

Errors to criterion. Analyses were performed on the number of each type of error (inhibition error, corresponding to a leverpress following the tone stimulus; omission error, corresponding to failure to leverpress following the light stimulus) made by the rats before reaching the criterion of performance at each ITI (not illustrated). A three-way ANOVA (lesion × type of errors × delay) performed on the total number of errors revealed no effect of lesion (F < 1). However, there was a significant effect of type of error [F(1,12) = 24.9, p < .001], indicating that the rats made significantly more inhibition errors than omission errors, as well as a significant effect of delay [F(3,36) = 9.8, p < .001], indicating better performance at the shorter ITIs than at the longer ones. None of the interactions between these three factors was significant (Fs < 1). Complementary analyses indicated a significant effect of the delay on the inhibition errors [F(3,36) = 34.38, p < .001], but not on the omission errors [F(3,36) = 2.45, p < .07].

Sessions to criterion. The number of training sessions necessary to reach the criterion at each ITI is illustrated in Figure 5. One rat in each group failed to reach all the criteria within 20 testing sessions. An ANOVA performed on these data did not reveal an effect of lesion (F < 1) but did indicate a significant effect of delay [F(3,36) = 14.61, p < .001], with more extensive training required for the longer ITIs. In addition, there was no lesion × delay interaction (F < 1), indicating that the lesions did not modulate the general effect of the delays.

GENERAL DISCUSSION

The present study confirmed that mPFC subregions in rats differentially contribute to cognitive function (Joel, Tarrasch, Feldon, & Weiner, 1997; Joel, Weiner, & Feldon, 1997; Kesner et al., 1996; Ragozzino et al., 1998; Ragozzino, Wilcox, et al., 1999; Seamans et al., 1995). In Experiment 1, lesions to the ACd cortex severely disrupted go/no-go response alternation performance, thereby replicating the impairment observed by Winocur (1991) in rats with more extensive lesions to the same area. In
contrast, lesions of the PL–IL area did not lead to a performance deficit on this task. This pattern of results was only partially expected. Although we predicted an impairment in response alternation rule learning in the ACd group, we also predicted, on the basis of previous evidence, performance deficits at relatively long delays in rats with PL–IL lesions (Brito & Brito, 1990; Delatour & Gisquet-Verrier, 1996, 1999, 2000).

To assess whether this discrepancy in results was due to different demands on the scheduling of responses, in Experiment 2, rats with PL–IL lesions were administered Delatour and Gisquet-Verrier’s (1999) conditional discrimination task, but with a significant change. In the original task, in which rats with PL–IL lesions were impaired at long ITIs, a block design was used in which ITIs were increased progressively over test sessions. In Experiment 2, the number and duration of ITIs were invariant throughout testing, as in Experiment 1 (see Winocur, 1991). Under these conditions, there was no difference between PL–IL and control groups. The finding that the same task may or may not be sensitive to PL–IL lesions, depending on the experimental procedure used (variable delays vs. block design), forces a reconsideration of the functional role of the PL–IL area. The implications of these results for understanding the respective roles of the dorsal and ventral mPFC regions are discussed in the following sections.

Functional Role for the Dorsal Anterior Cingulate Region

As was noted by Winocur (1991), the impairment of ACd rats in the VI delayed alternation task reflects a deficit in learning the alternation rule that is independent of the ITI. It is important to emphasize that ACd lesions did not obliterate performance on this task, since all the rats exhibited significant improvement throughout training. Nevertheless, relative to PL–IL and control rats, the ACd group exhibited widespread impairment that was reflected primarily in a failure to modulate their response latencies according to the reinforcement schedule. This effect was particularly clear for the no-go trials, where the response latencies of ACd rats were substantially shorter than those of the other two groups.

There has been relatively little investigation of the behavioral consequences of lesions restricted to the ACd region. The few available studies report little or no effect of these lesions on acquisition processes. For example, ACd lesions did not affect performance on tasks in which rats were required to discriminate arms of a maze on the basis of spatial or visual cues (Ragozzino et al., 1998; Ragozzino, Wilcox, et al., 1999). Rats with ACd lesions were found to be slightly or not impaired on various spatial learning and memory tasks (e.g., Joel, Tarrasch, et al., 1997; Joel, Weiner, & Feldon, 1997; Sánchez-Santed, de Bruin, Heinsbroek, & Verwer, 1997; Seamans et al., 1995),
but interestingly, when obtained, these effects were always independent of working memory load. These observations are consistent with the results of Experiment 1, in which the effects of ACd lesions did not interact with the effects of delay, as would have been expected if working memory was a factor in this group's deficit.

The main difficulties resulting from ACd lesions in Experiment 1 of the present research were associated with an inability to withhold leverpressing during the no-go trials. This pattern suggests a deficit in inhibitory processes, resulting in an uncontrolled tendency to emit a well-learned, reinforced response. Such a hypothesis would account for the perseverative tendencies that have been observed in ACd-lesioned rats in other tasks (Sánchez-Santed et al., 1997; Seamans et al., 1995). However, arguing against this view is the finding that ACd-lesioned rats were able to withhold inappropriate responding in a go/no-go spatial recognition test and in a simple spatial discrimination task (Ragozzino et al., 1998).

Another hypothesis that would account for the learning impairment of the alternation rule in Experiment 1 is that ACd lesions interfered with response selection. A deficit in response selection has been associated with large dorsal mPFC lesions involving both the ACd and the PrCm cortices, in various conditional learning tasks (Passingham et al., 1988; Winocur, 1992; Winocur & Eskes, 1998). In these tasks, lesioned rats were consistently impaired in selecting correct responses from various alternatives. However, when lesions are restricted to the ACd, deficits in response selection are not reliably obtained. In Experiment 1 of the present research, response selection deficits were observed only in the no-go trials, whereas a hypothesis that emphasizes this type of deficit would have predicted poor response selection on go and no-go trials. Furthermore, ACd lesions did not impair performance when rats were required to switch from a spatial to a visual-cued version of a discrimination task, suggesting that these lesions do not interfere with response selection strategies (Ragozzino, Wilcox, et al., 1999). Finally, we recently observed that rats with ACd lesions performed normally on a conditional discrimination task that required response selection on the basis of correct stimulus discrimination (Delatour & Gisquet-Verrier, 1997; see also Jackson, Kesner, & Aman, 1998, for similar results following dorsal mPFC lesions involving the ACd). Taken together, these results indicate that lesions restricted to the ACd preserve response selection in a diverse set of circumstances.

A comparison of tasks in which rats with ACd lesions exhibit normal response selection and those in which they do not may provide the basis of an alternate hypothesis of the functional role of this region. The above-mentioned tasks that are insensitive to the effects of such lesions provide a clear choice of response alternatives that is governed by salient discriminative stimuli (see also Joel, Tarrasch, et al., 1997; Joel, Weiner, & Feldon, 1997). An important feature of these tasks is that they do not require temporal organization or response sequencing. By comparison, successful performance on the go/no-go alternation task requires that rats pattern their responses according to different reinforcement contingencies that co-exist with variable ITIs. To accomplish this, rats must adopt a strategy that links conditional rule learning, the processing of temporally ordered information, and response selection demands. We hypothesize that the specific role of the ACd is in the implementation of rule-based behavior that requires the integration of temporal cues and response sequencing components. This view derives support from earlier evidence that rats with dorsal mPFC lesions involving the ACd are impaired on various learning tasks that challenge timing ability (Chaillan, Marchetti, Delfosse, Roman, & Soumireu-Mourat, 1997; Dietrich & Allen, 1998; Olton, 1989), require temporal ordering of responses (Chiba, Kesner, & Reynolds, 1994; Kesner & Holbrook, 1987; Mogensen & Divac, 1984; Mogensen & Holm, 1994), or engage memory for frequency (Kesner, 1990).

**Functional Role for the Prelimbic-Infralimbic Area**

The results of Experiments 1 and 2 showed that PL–IL lesions did not affect rats' performance on our VI, delayed alternation, and delayed conditional discrimination learning tasks. This finding was somewhat surprising, given the severe delay-dependent deficits of rats with PL–IL lesions on a variant of the conditional discrimination learning task (Delatour & Gisquet-Verrier, 1999). However, there was a major difference between the tasks, in that the latter involved the progressive introduction of delays between stimulus presentation and the opportunity to respond, whereas, in the former, the same critical delays were always present throughout testing. Thus, it would appear that time-dependent deficits following PL–IL lesions are related to the way that delays are introduced during testing. It is clear from the results that PL–IL lesions did not affect the ability to retain specific information. Rather, they interfered with the capacity to react adequately to a longer delay that represents a change in the experimental situation, requiring behavioral adjustment. This contention is supported by previous results (Delatour & Gisquet-Verrier, 1999) showing that extending the delays, in a block design, constantly increased the number of errors made by rats with PL–IL lesions. The same effect was observed to a lesser degree in control rats. In Experiment 2 of the present research, the control group received 56 trials/sessions on the variable-delay procedure and reached criterion in about 16 sessions. By comparison, Delatour and Gisquet-Verrier (1999) administered 70 trials/session in the progressive-delay procedure, and control rats required 22 trials to learn the task. Thus, a progressive-delay procedure increases task difficulty and appears to force a reengagement of learning processes that facilitates behavioral adjustment to the new task contingencies. The importance of this variable with respect to PL–IL function is highlighted by...
other evidence that delay-dependent deficits following lesions to this area are consistently associated with the progressive-delay procedure (Brito & Brito, 1990; Delatour & Gisquet-Verrier, 1996, 1999, 2000). In contrast, such deficits have not been reported in tasks that employed a variable-delay procedure (Aggleton, Neave, Nagle, & Sahgal, 1995; Chudasama & Muir, 1997; Ragozzino et al., 1998).

The present results showing no effect of PL–IL lesions even for extended delays is of particular importance because they question the hypothesis that this area is involved in working memory (Brito & Brito, 1990; Delatour & Gisquet-Verrier, 1996, 1999, 2000; Ragozzino et al., 1998). Kesner and his colleagues (Kesner, 2000; Ragozzino et al., 1998) take the view that involvement of the PL–IL region in working memory is restricted to spatial and visual object information. Although this hypothesis could be applied to the present results, we have already noted that a working memory hypothesis, even one that is restricted to the spatial modality, cannot fully account for the behavioral deficits resulting from damage to this area. It is well established that PL–IL lesions do not consistently affect performance in spatial working memory tasks. For example, rats with PL–IL lesions performed normally in a standard elimination task in a radial arm maze (Delatour & Gisquet-Verrier, 1996), even when working memory was challenged by introducing delays between arm entries (Joel, Tarrasch, et al., 1997). As well, it was found that PL–IL lesions interfered with performance on spatial reference memory tasks that required a high level of behavioral flexibility but in which there were no working memory components (Delatour & Gisquet-Verrier, 2000; see also Granon & Poucet, 1995, for similar results).

An alternative hypothesis that takes into account the various effects of PL–IL lesions emphasizes the structure’s role in processes related to attention and behavioral flexibility. PL–IL involvement in attentional mechanisms has been suggested by other investigators (Bussey et al., 1997; Granon, Hardouin, Courtier, & Poucet, 1998; Granon, Save, & Buhot, 1996), following observations of lesion-induced deficits in several experimental paradigms. For example, Granon and colleagues (Granon et al., 1998) showed that the PL cortex plays a central role in maintaining sustained attention that is necessary to detect subtle brightness variations. As well, there is growing evidence that the PL–IL area contributes to behavioral flexibility. For example, de Bruin, Sánchez-Santed, Heinsbroek, Donker, and Postmes (1994) reported that PL–IL lesions disrupted rats’ ability to switch from a spatial to a cued version of the hidden-platform task in the Morris water maze. Other investigators have demonstrated similar switching difficulties in other paradigms in rats subjected to temporary or permanent disruption of PL–IL function (Joel, Weiner, & Feldon, 1997; Ragozzino, Detrick, & Kesner, 1999; Ragozzino, Wilcox, et al., 1999; Seamans et al., 1995). As well, there is evidence that PL–IL lesions do not interfere with some forms of behavioral flexibility, as is required, for example, in reversal learning. For instance, PL–IL rats were able to adapt their performance in a task involving four baited arms and four nonbaited arms in a radial maze when the two sets of arms were reversed (Joel, Tarrasch, et al., 1997) and in a water maze when the location of the hidden platform was modified (Granon & Poucet, 1995). The lack of effect of PL–IL lesions on reversal learning may be modulated by task difficulty (Bussey et al., 1997). These investigators found that mPFC lesions (centered around the PL area) did not affect reversal learning for easily discriminable stimuli (stimuli that differed in size and brightness), but they disrupted reversal for stimuli that were more difficult to discriminate (stimuli that only differed in brightness).

The hypothesis that behavioral deficits associated with PL–IL damage can be linked to demands on cognitive processes that control adjustments to changes in task contingencies is consistent with a large body of evidence. However, the relative involvement of attentional and response control mechanisms needs to be clarified. In this regard, a recent study by Ragozzino, Detrick, and Kesner (1999) is particularly interesting. These investigators trained rats in a place or in a response-learning task in a cross-maze and then proposed either a reversal of the initial discrimination or a switch of the discrimination rule. They found that PL–IL inactivation affected behavioral flexibility only for cross-modal switching (spatial vs. cued version), but not for intramodal switching (reversal learning), supporting the view that the PL–IL area is involved in both attention and behavioral flexibility.

Conclusion

The present results strengthen the contention that the subregions of the mPFC in rats contribute differentially to behavior and, in addition, provide insight into the respective functions of the ACd and the PL–IL regions. The results, in line with those of other laboratories, support the hypotheses that the ACd region is involved in sequencing of temporally ordered behavior, whereas the PL–IL area is critically involved in processes related to attentional mechanisms and behavioral flexibility.

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