Medial prefrontal cortex is selectively involved in response selection using visual context in the background

Inah Lee¹ and Ji Yun Shin
Department of Brain and Cognitive Sciences, Seoul National University, Gwanak-ro, Gwanak-gu, Seoul 151-742, Korea

The exact roles of the medial prefrontal cortex (mPFC) in conditional choice behavior are unknown and a visual contextual response selection task was used for examining the issue. Inactivation of the mPFC severely disrupted performance in the task. mPFC inactivations, however, did not disrupt the capability of perceptual discrimination for visual stimuli. Normal response selection was also observed when nonvisual cues were used as conditional stimuli. The results strongly suggest that the mPFC is not necessarily involved in the inhibition of response or flexible response selection in general, but is rather critical when response selection is required conditionally using visual context in the background.

"Contextual response selection" enables an organism to respond flexibly and adaptively in various situations when encountering similar objects (potentially associated with conflicting responses) across different contextual settings. It is implicated in the literature that the prefrontal cortex (PFC) is critically involved in making flexible responses across different contexts (Kesner and Ragozzino 2003; Haddon and Killcross 2006; Marquis et al. 2007; Lee and Solivan 2008; Horga et al. 2011). The PFC receives direct afferent fibers from the hippocampus (Jay and Witter 1991; Thierry et al. 2000), which is one of the critical structures for contextual memory tasks (Hirsh 1974; Good and Honey 1991; Kim andFanselow 1992; Kim and Lee 2011). Presumably related to such "contextual connections," perturbations of PFC lead to severe impairment in contextual response selection tasks in which visual contextual cues play significant roles (Haddon and Killcross 2006; Jo et al. 2007; Marquis et al. 2007; Lee and Solivan 2008).

In prior studies, however, behavioral tasks required subjects to learn complicated multiple relationships between noncontextual stimuli (e.g., object, light, tone, etc.) and visual contexts (e.g., room cues) when testing contextual response selection (Haddon and Killcross 2006; Marquis et al. 2007; Lee and Solivan 2008). In other cases, rats learned biconditional associations between two different objects and locations in a maze and flexibly responded to a context-relevant object for obtaining reward (Lee and Solivan 2008). What these studies have tested is whether rats could learn associations between multiple individual cues and discrete contexts and whether PFC was necessary in flexibly responding (or not responding) to particular cue-context paired associates. It is unclear in these behavioral paradigms, however, how much “response selection” component was tested and also what proportion of resulting performance deficits following PFC manipulations could be attributed to contextual response selection per se. In the current study, in order to establish a more straightforward relationship between context and response selection behavior, we removed the requirement of associating other sensory cues (e.g., light, tone, and objects) with context and just made a single object (i.e., sand-filled jar) allow two distinctively different types of motor responses (digging the sand in the jar or pushing the jar).

Rats (n = 8, male Long-Evans) were trained in a visual contextual response selection (VCRS) task (Fig. 1A,B) and were implanted with bilateral cannulae (26G) targeting the mPFC (2.7 mm anterior to bregma, 1.5 mm lateral to midline at a 10° angle, 4.9 mm ventral to the skull surface) (Fig. 1C) once they reached performance criterion (≥75% correct choices for both contexts for two consecutive days). In the VCRS task (32 trials/session), the rat was required to push a sand-filled jar (480 g) when an array of peripheral LCD panels (each panel, 17” TFT LCD monitor) displayed a certain visual context (e.g., pebbles pattern) (Fig. 1A), but the rat must dig the sand in the same object when another visual context (e.g., zebra pattern) (Fig. 1B) was shown in order to obtain the reward (cereal). The stimulus–response relationships associated with rewards were counterbalanced among the rats. The jar was placed in the center of the surrounding LCD panels in the background. We prevented rats from using a potential olfactory strategy by mixing sand with ground Cheerios powder (1:5 ratio). Rats were unable to tell whether a piece of cereal was in the sand or not by smelling it because they showed digging behavior even when the reward was missing in some probe trials. Since the jar was heavy and the cereal reward was buried deep in the jar, the two response modes were equally effortful behaviors in which the rat should be intentionally engaged for obtaining reward, and thus were easily discriminated by the experimenter in the testing room. If the rat moved the jar (even a little without exposing the food well underneath it), it was considered a pushing response. Once the rat touched the sand with both front paws, it was considered a digging response. No correction was allowed once a wrong behavioral selection was made. One of the rats was dropped from further analysis due to misplacement of cannula tips. For the postsurgical testing of performance, phosphate-buffered saline (SAL, 0.5 μL per site) was injected on day 1 and muscimol (MUS, 0.5 μg/0.5 μL per site), a GABA-A receptor agonist, was injected afterward for 2 d in a row. SAL was injected again on the last day for checking the baseline performance. All cannula tip positions of the remaining seven rats were located in the mPFC (Fig. 1C). The diffusion range of MUS was estimated using fluorescent MUS after all experiments were finished (see Jo and Lee 2010 for detailed methods) and the results showed that MUS diffusion covered mostly the prelimbic and infralimbic cortices (spanning anteroposteriorly +3.5 to +1.9 mm from bregma and ±1.2 mm mediolaterally) on average. Rats injected with SAL in the mPFC before and after MUS injections were normal in performing the
VCRS task (Fig. 1D). The same rats, however, were severely impaired when MUS was injected in the mPFC compared with SAL conditions (Fig. 1D). A repeated-measures ANOVA showed a highly significant effect of drug condition \( (F_{1,3,21} = 20.3, P < 0.0001) \). Post hoc comparisons (Tukey-Kramer) revealed significant differences between all SAL and MUS conditions \( (P\text{-values of} < 0.05) \). However, despite the seemingly increased performance in the second MUS condition (Fig. 1D), no statistical significance was found between the first and second MUS conditions. No significant difference in performance was found between SAL conditions. No significant effect of drug was found when average response latency (from start-box exit to response) was compared across days, suggesting that MUS did not disrupt generic sensory-motor capabilities. The results strongly demonstrate that the mPFC is critical for visual contextual response selection.

In the VCRS task, the rat must first identify the surrounding visual context presented in a given trial in order to choose a correct response associated with the context. We examined whether the capability of visual discrimination was impaired in the mPFC-inactivated rats using a T-maze in a visual cue discrimination (VCD) task (Fig. 2A). In the VCD task (32 trials/session), the rat was simply required to turn to one of the arms associated with the peripheral LCD monitor displaying the rewarding visual stimulus and to displace a metal washer to retrieve a cereal reward (Fig. 2A). Once a trial started, the rat exited the start box and entered the response platform. The animal was able to sample the tactile cue on the floor as it entered the platform until it responded to the sand-filled jar either with a digging or pushing response (depending on the tactile cue). The relationships between tactile stimuli and dig–push responses were counterbalanced among rats. The same within-subject drug-injection schedule (SAL-MUS-MUS-SAL) used for prior tasks was used. The rats injected with MUS in the mPFC performed as well (>80%) as in SAL conditions and showed normal tactile-cued responses to the sand-filled jar (Fig. 3B). No significant drug effect was observed \( (F_{1,3,21} = 2.75, n.s.; \text{repeated-measures ANOVA}) \).

One may reason that the null effects of MUS in the mPFC in both VCD and TCRS tasks might be related to a possibility that MUS could have exerted less inhibitory influence on the mPFC in later days of testing as the system was exposed to MUS for multiple days throughout the study. This is unlikely because, when tested in the VCRS task after the completion of the TCRS task, MUS again severely disrupted performance in the same animals, whereas no such impairment was observed with SAL (Fig. 3C) \( (F_{1,7} = 15.39, P < 0.01, \text{repeated-measures ANOVA}) \). The results overall strongly suggest that mPFC is selectively involved in contextual response selection involving distal visual cues in the background.

The functional involvement of mPFC in response selection was apparently limited to the VCRS task in which the contextual stimulus served as a critical conditional cue implicitly from the background at the time of conditional response selection. The background visual stimuli in the VCRS task worked as contextual cues implicitly or indirectly at the time of response selection as compared with the explicit elemental cues in both VCD and TCRS tasks. Hirsh (1974) emphasized this implicit nature of influence by background stimuli when defining contextual learning and memory in the hippocampus. Although the current study manipulated mPFC instead of the hippocampus, the results connect well to such contextual theory. Alternatively, one may focus on the difference between distal and local cues to explain different results between VCRS and TCRS tasks. However, the mPFC-inactivated rats were normal in the VCD task where distal cues were also used as in the VCRS task. Therefore, the impairment in the VCRS
task may not be simply attributable to the distal nature of visual stimulus. Despite the difference in the type of behavioral responses between VCRS and VCD tasks (i.e., different behavioral interactions with the same object vs. simple orienting response toward a visual cue), complexity in motor response per se may not be a critical factor underlying the performance deficits with mPFC inactivations, because the TCRS task was identical to the VCRS task with respect to the required behavioral responses. The results from the TCRS task strongly suggest that some of the key cognitive components (including resolving conflicts between different behavioral responses) for conditionally choosing a proper response toward the same object were intact without the mPFC. Literature also suggests that mPFC-inactivated rats are normal in inhibiting inappropriate responses and flexibly choosing correct ones in certain situations (Bussey et al. 1997; Delatour and Gisquet-Verrier 1999; Chudasama et al. 2003; Ragozzino et al. 2003; McDonald et al. 2007; Floresco et al. 2008; Hayton et al. 2010).

Attention related to task difficulty may also explain the differential influences of MUS in the mPFC in different tasks in our study. Specifically, several hypotheses have been proposed for explaining mPFC functions, including working memory (Goldman-Rakic 1990; Granon et al. 1994; Delatour and Gisquet-Verrier 1996), response selection (Delatour and Gisquet-Verrier 1996; Haddon and Killcross 2006; Marquis et al. 2007), high-level attention or effortful processing (Granon et al. 1995; Delatour and Gisquet-Verrier 1996; Muir et al. 1996; Bussey et al. 1997), response inhibition (Chudasama et al. 2003; McDonald et al. 2007; Hayton et al. 2010), and flexible task/rule-switching (Ragozzino et al. 2003; Floresco et al. 2008; Oualian and Gisquet-Verrier 2010), to name a few. Among these hypotheses, the results from the current study may be explained most parsimoniously by the attention-based hypothesis; that is, the mPFC plays significant roles when higher-level attention is required for goal-directed response in a task (Granon and Poucet 1995; Granon et al. 1998).

Figure 2. Visual cue discrimination (VCD) task. (A) Behavioral paradigm. The rat was rewarded when displacing the washer in the arm in front of the LCD monitor that showed a rewarding visual stimulus. (B) Performance in the VCD task with SAL and MUS infusions in the mPFC. No significant impairment was observed in performance with mPFC inactivation compared with SAL conditions. (Mean ± SEM.)

Figure 3. Tactile-cued response selection (TCRS) task (A, B) and VCRS task (C). (A) Behavioral paradigm. A floor insert with tactile cue (left, soft shelf liner; right, wire mesh) was used for informing which response (pushing or digging) was appropriate toward the same object (sand-filled jar) for obtaining a reward in a given trial. (B) Rats showed no significant differences in performance between SAL and MUS conditions. (Mean ± SEM.) (C) Performance in the VCRS task run after the TCRS task in the same rats. The rats remembered the VCRS task well with SAL injections in the mPFC, whereas performance was severely affected again by MUS injections. (Mean ± SEM.)
showed that rats with mPFC lesions were not impaired in a conditional task in a Y-maze using simple visual stimulus (e.g., right arm for slowly flashing light and left arm for fast flashing light). Supporting this line of reasoning, it took 2–3 wk for the rats to learn the VCRS task, whereas it took only 4 d on average for the rats to learn the VCD task and approximately the same amount of time for the TCRS task, suggesting that the latter two tasks were relatively easier than the VCRS task. The attention-based explanation, however, needs to be further examined with other tasks with matching attention levels (but without using visual stimuli) in the future.

In sum, the current study demonstrates the importance of the mPFC when conditional response should be made using visual context in the background to an otherwise ambiguous object. However, flexible behavioral selection was possible without normal function of the mPFC when the choice behavior was cued by elemental, but not contextual cues. The results should provide important clues for future studies to further delineate conditions in which the mPFC plays critical roles in choice behavior.

Acknowledgments

The current study was supported by the World Class University program of the Korea Science and Engineering Foundation funded by the Ministry of Education, Science and Technology (R31-10089), as well as by NIMH RO1 MH079971.

References

Bussey TJ, Muir JL, Everitt BJ, Robbins TW. 1997. Triple dissociation of anterior cingulate, posterior cingulate, and medial frontal cortices on visual discrimination tasks using a touchscreen testing procedure for the rat. Behav Neurosci 111: 920–936.

Chudasama Y, Passetti F, Rhodes SE, Lopian D, Desai A, Robbins TW. 2003. Dissociable aspects of performance on the 5-choice serial reaction time task following lesions of the dorsal anterior cingulate, infralimbic and orbitofrontal cortex in the rat: Differential effects on selectivity, impulsivity and compulsivity. Behav Brain Res 146: 105–119.

DeCoteau WE, McElvaine D, Smolentzov L, Kesner RP. 2009. Effects of rodent prefrontal lesions on object-based, visual scene memory. Neurobiol Learn Mem 92: 552–558.

Delatour B, Gisquet-Verrier P. 1996. Prelimbic cortex specific lesions disrupt delayed-variable response tasks in the rat. Behav Neurosci 110: 1282–1298.

Delatour B, Gisquet-Verrier P. 1999. Lesions of the prelimbic-infralimbic cortices in rats do not disrupt response selection processes but induce delay-dependent deficits: Evidence for a role in working memory? Behav Neurosci 113: 941–955.

Floresco SB, Block AE, Tse MTL. 2008. Inactivation of the medial prefrontal cortex of the rat impairs strategy set-shifting, but not reversal learning, using a novel, automated procedure. Behav Brain Res 190: 85–96.

Goldman-Rakic PS. 1990. Cellular and circuit basis of working memory in prefrontal cortex of nonhuman primates. Prog Brain Res 85: 325–335.

Good M, Honey RC. 1991. Conditioning and contextual retrieval in hippocampal rats. Behav Neurosci 105: 499–509.

Granon S, Pouget B. 1995. Medial prefrontal lesions in the rat and spatial navigation: Evidence for impaired planning. Behav Neurosci 109: 474–484.

Granon S, Vidal C, Thinus-Blanc C, Changeux JP, Pouget B. 1994. Working memory, response selection, and effortful processing in rats with medial prefrontal lesions. Behav Neurosci 108: 883–891.

Granon S, Pouget B, Thinus-Blanc C, Changeux JP, Vidal C. 1995. Nicotinic and muscarinic receptors in the rat prefrontal cortex: Differential roles in working memory, response selection and effortful processing. Psychopharmacology (Berl) 119: 139–144.

Granon S, Save E, Buhot MC, Pouget B. 1996. Effortful information processing in a spontaneous spatial situation by rats with medial prefrontal lesions. Behav Brain Res 78: 147–154.

Granon S, Hardouin J, Courtier A, Pouget B. 1998. Evidence for the involvement of the rat prefrontal cortex in sustained attention. Q J Exp Psychol B 51: 219–233.

Haddon JE, Killcross S. 2006. Prefrontal cortex lesions disrupt the contextual control of response conflict. J Neurosci 26: 2933–2940.

Hayton SJ, Lovett-Barron M, Dumont EC, Olstead MC. 2010. Target-specific encoding of response inhibition: increased contribution of AMPA to NMDA receptors at excitatory synapses in the prefrontal cortex. J Neurosci 30: 11493–11500.

Harsh R. 1974. The hippocampus and contextual retrieval of information from memory: A theory. Behav Biol 12: 421–444.

Horga G, Maia TV, Wang P, Wang Z, Marsh R, Peterson BS. 2011. Adaptation to conflict via context-driven anticipatory signals in the dorsomedial prefrontal cortex. J Neurosci 31: 16208–16216.

Jay TM, Witter MP. 1991. Distribution of hippocampal CA1 and subicular efferents in the prefrontal cortex of the rat studied by means of anterograde transport of Phaseolus vulgaris-leucoagglutinin. J Comp Neurol 315: 574–586.

Jo YS, Lee I. 2010. Disconnection of the hippocampal–perihitnial cortical circuits severely disrupts object–place paired associative memory. J Neurosci 30: 9850–9858.

Jo YS, Park EH, Kim IH, Park SK, Kim H, Kim HT, Choi JS. 2007. The medial prefrontal cortex is involved in spatial memory retrieval under partial-cue conditions. J Neurosci 27: 13567–13578.

Kesner RP, Ragozzino ME. 2003. The role of the prefrontal cortex in object-place learning: A test of the attribute specificity model. Behav Brain Res 146: 159–165.

Kim JJ, Fanselow MS. 1992. Modality-specific retrograde amnesia of fear. Science 256: 675–677.

Kim J, Lee I. 2011. Hippocampus is necessary for spatial discrimination using distal cue-configuration. Hippocampus 21: 609–621.

Lee I, Solivan F. 2008. The roles of the medial prefrontal cortex and hippocampus in a spatial paired-association task. Learn Mem 15: 357–367.

Marquis J, Killcross S, Haddon JE. 2007. Inactivation of the prelilmbic, but not infralimbic, prefrontal cortex impairs the contextual control of response conflict in rats. Eur J Neurosci 25: 559–566.

McDonald R, Foong N, Ray C, Rizos Z, Hong N. 2007. The role of medial prefrontal cortex in context-specific inhibition during reversal learning of a visual discrimination. Exp Brain Res 177: 509–519.

Muir JL, Everitt BJ, Robbins TW. 1996. The cerebral cortex of the rat and visual attentional function: Dissociable effects of mediofrontal, cingulate, anterior dorsolateral, and parietal cortex lesions on a five-choice serial reaction time task. J Comp Neurol 361: 251–262.

Olual C, Gisquet-Verrier P. 2010. The differential involvement of the prelilmbic and infralimbic cortices in response conflict affects behavioral flexibility in rats trained in a new automated strategy-switching task. Learn Mem 17: 654–668.

Ragozzino ME, Kim J, Hassert D, Minniti N, Kiang C. 2003. The contribution of the rat prelilmbic-infralimbic areas to different forms of task switching. Behav Neurosci 117: 1054–1065.

Richier A-M, Gioanni Y, DeGenetais E, Glowinski J. 2000. Hippocampoprefrontal cortex pathway: anatomical and electrophysiological characteristics. Hippocampus 10: 411–419.

Winocur G, Eskes G. 1998. Prefrontal cortex and caudate nucleus in conditional associative learning: Dissociated effects of selective brain lesions in rats. Behav Neurosci 112: 89–101.

Received February 7, 2012; accepted in revised form April 12, 2012.