The contribution of cholinergic and dopaminergic afferents in the rat prefrontal cortex to learning, memory, and attention

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Recent experiments have provided valuable information about the functional organization of the subregions of the rat prefrontal cortex. This review assesses the findings from studies in which the role of the cholinergic and dopaminergic systems in the anterior cingulate, prelimbic, and infralimbic areas was investigated by using several different approaches—that is, chemical lesions, pharmacological manipulations, or in vivo microdialysis. Cholinergic and dopaminergic input to the prelimbic and infralimbic subregions may modulate multiple behavioral processes. Activation of the muscarinic cholinergic receptors or activation of the dopamine D1 receptors in the prelimbic and infralimbic areas modulates working memory processes. Cholinergic input to the prelimbic and infralimbic areas may also modulate attention, since changes in acetylcholine release occur when there is an increase in attentional demands. In addition, activation of dopaminergic receptors in the prelimbic and infralimbic areas influences behavioral flexibility. Cholinergic input to the anterior cingulate does not appear critical for working memory but likely contributes to attentional processes. Overall, the results suggest that acetylcholine and dopamine affect multiple behavioral functions subserved by the rat prefrontal cortex that depend on modulating activity in specific prefrontal cortex subregions.

The human prefrontal cortex is thought to be a region of the cerebral cortex that is intimately involved in some of the most complex behavioral and cognitive functions (Fuster, 1997; Shallice & Burgess, 1991; Stuss & Benson, 1986). Neuropsychological assessments of prefrontal-damaged patients commonly reveal dysfunctions in attention, working memory, strategy selection, and temporal sequencing (Burgess & Shallice, 1996; Chao & Knight, 1995; Damasio, Tranel, & Damasio, 1990; Freedman, Black, Ebert, & Binns, 1998; Kesner, Hopkins, & Fineman, 1994; McDowell, Whyte, & DeEsposito, 1998; Milner, Petrides, & Smith, 1985; Owen, 1997; Owen, Morris, Sahakian, Polkey, & Robbins, 1996; Swain, Polkey, Bullock, & Morris, 1998). Largely because the human prefrontal cortex is proposed to be a brain region involved in high-level cognitive processes, attempts to understand the neural mechanisms underlying these processes have involved experiments carried out predominantly in non-human primates. Nevertheless, all mammalian species appear to have a prefrontal cortex, although there is still debate on how to anatomically define the prefrontal cortex in some species (Preuss, 1995; Uylings & van Eden, 1990). In addition, during recent decades, there have been numerous experiments in rodents that suggest that this species can serve as a useful model for understanding prefrontal cortex functioning (Kolb, 1984, 1990). In particular, pharmacological manipulations and lesions of the rat prefrontal cortex lead to behavioral deficits comparable with those found in humans with prefrontal cortex damage—that is, impairments in working memory, strategy selection, and temporal sequencing (Brito & Brito, 1990; de Bruin, Swinkels, & De Brabander, 1997; DeCoteau, Kesner, & Williams, 1997; Frysztak & Neafsey, 1994; Gallagher, McMahan, & Schoenbaum, 1999; Granon & Pouet, 1995; Kesner & Holbrook, 1987; Kesner, Hunt, Williams, & Long, 1996; Kolb, Burham, McDonald, & Sutherland, 1994; Muir, Everitt, & Robbins, 1996; Ragozzino, Adams, & Kesner, 1998; Ragozzino, Detrick, & Kesner, 1999a; Ragozzino & Kesner, 1998; Ragozzino, Wilcox, Raso, & Kesner, 1999).

Another similarity in the experimental findings among different mammalian species is that separate prefrontal cortex subregions differentially contribute to various behavioral and cognitive functions (Bussey, Muir, Everitt, & Robbins, 1997; DeCoteau et al., 1997; Dias, Robbins, & Roberts, 1996; Eichenbaum, Clegg, & Feeley, 1983; Fuster, 1997; Kesner et al., 1996; Mogensen & Holm, 1994; Petrides, Alivisatos, Evans, & Meyer, 1993; Ragozzino et al., 1998; Ragozzino, Detrick, & Kesner, 1999a; Seamans, Floresco, & Phillips, 1995; Wilson, Scalaidhe, & Goldman-Rakic, 1993). However, an understanding of the mechanisms in different prefrontal cortex subregions that underlie various functions remains incomplete. In recent years, there has been an interest in determining the neurochemical changes in different parts of the prefrontal cortex that influence various behavioral processes—that
is, learning and attention. These experiments have involved the use of different techniques, such as intracranial drug injections, specific neurotoxic lesions, and in vivo microdialysis. Although the rat prefrontal cortex consists of a medial and a lateral sector, the vast majority of these studies have concentrated on the neurochemical mechanisms in the medial prefrontal cortex—particularly the anterior cingulate, prelimbic, and infralimbic subregions—that underlie particular cognitive processes. This review focuses on cholinergic and dopaminergic actions in the rat anterior cingulate, prelimbic, and infralimbic subregions as they relate to different behavioral processes involving attention, learning, and memory. The discussion will be limited to studies directly measuring or manipulating dopamine and acetylcholine changes in these prefrontal cortex subregions. In some of the experiments discussed, the role of these neurochemicals in only the prefrontal region was examined, whereas in other investigations, their role in the prefrontal and infralimbic areas combined was examined. These findings are discussed under one heading, but whether a particular study involved only the prefrontal region or both the prefrontal and the infralimbic areas is duly noted.

ANATOMY

The rat prefrontal cortex is a heterogenous structure that consists of anatomically distinct subregions that may also represent separate functional compartments. The prefrontal cortex is commonly defined by either reciprocal or comparatively dense connections with the mediodorsal thalamic nucleus, relative to other thalamic nuclei (Groenewegen, 1988; Uylings & van Eden, 1990). The connections between the mediodorsal thalamic nucleus and the prefrontal cortex are topographically organized, leading to the following two subdivisions of the rat prefrontal cortex: (1) a medial division consisting of the medial precentral area, the dorsal and ventral anterior cingulate, and prefrontal, infralimbic, and medial orbital areas and (2) a lateral division encompassing the lateral orbital cortex and the dorsal and ventral agranular insular area (Groenewegen, 1988; Krettek & Price, 1977; Leonard, 1969; Uylings & van Eden, 1990).

The main source of cholinergic afferents to the rat prefrontal cortex is the basal forebrain (Gaykema, Luiten, Nyakas, & Traber, 1990; Woolf, 1991). In the basal forebrain, the nucleus basalis magnocellularis and the horizontal limb of the diagonal band area are the principal regions that send cholinergic projections to the prefrontal cortex (Gaykema et al., 1990; Woolf, 1991). The vertical diagonal band and medial septal area appear to have a sparse cholinergic projection to the prefrontal cortex (Gaykema et al., 1990). Furthermore, the laterodorsal tegmental nucleus also has cholinergic projections to the ventromedial prefrontal cortex (Satoh & Fibiger, 1986).

The dopaminergic afferents to the rat prefrontal cortex originate from the midbrain tegmentum (Berger, Gaspar, & Verney, 1991). The dopamine terminals in the ventral portions of the medial prefrontal cortex originate from the ventral tegmental area and terminate predominantly in layers V and VI (Berger et al., 1991). The dopamine projections to the anterior cingulate originate mainly from the substantia nigra and terminate in the superficial layers (Berger et al., 1991).

THE ROLE OF CHOLINERGIC TRANSMISSION IN THE PREFRONTAL CORTEX ON COGNITIVE PROCESSES

Most experiments that have focused on how cholinergic transmission in the medial prefrontal cortex influences behavioral and cognitive processes have employed working memory and attention tasks. In addition to traditional pharmacological approaches, in vivo microdialysis and the selective cholinergic neurotoxin 192 IgG-saporin are now being employed to facilitate an understanding of how cholinergic transmission in separate brain regions influences different behavioral and cognitive processes. This section will include some of the most recent experiments that have used these techniques.

Cholinergic Influences in the Prelimbic and Infralimbic Areas on Working Memory

The findings from several studies suggest that the rat prefrontal cortex, like the nonhuman primate and human prefrontal cortices, is involved in working memory (DeCoteau et al., 1997; Delatour & Gisquet-Verrier, 1996; Granon, Poucet, Thinas-Blanc, Changeux, & Vidal, 1995; Kesner et al., 1996; Kolb et al., 1994; Ragozzino et al., 1998; Seamans et al., 1995). Working memory commonly refers to the short-term storage of information that enables accurate execution of a response. Spatial delayed alternation and delayed nonmatching-to-sample tasks are frequently used to assess working memory processes. In the spatial delayed alternation task, a rat is often tested in a T-maze in which its choice of which arm to enter is always between the same two arms. A rat is reinforced for choosing either arm on the first trial, and on each subsequent trial, it must alternate and always choose the opposite arm to that chosen on the previous trial to receive a reinforcement (or avoid an aversive stimulus—i.e., a shock). A delay is inserted between trials, during which the rat must remember which arm it had just entered on a trial.

In contrast to the spatial delayed alternation task, each trial in a delayed nonmatching-to-sample task is usually divided into a sample phase and a test phase. In the sample phase, a stimulus is presented that the subject must respond to. After a delay, the test phase begins, in which two stimuli are presented—one that is identical to the sample phase and one that is novel for that trial. The novel stimulus must be chosen to receive a reinforcement. In order to choose the novel stimulus, the stimulus that was presented during the sample phase has to be remembered.

Although the spatial delayed alternation and the delayed nonmatching-to-sample tasks have a working mem-
ory component to them, these tasks also take advantage of rats' natural tendency to alternate (Dember & Fowler, 1958; Olton & Schlosberg, 1978). Therefore, the tasks also have a strategic component to them, in which rats have a bias to choose the location that has been least recently visited or the novel stimulus. This raises the possibility that a deficit in one of these tasks may result from an impairment in proper execution of an alternation strategy, as opposed to a working memory deficit. If a deficit in these tasks is due to a selective impairment in using a strategy, the deficit should occur independent of the delay. However, if a particular manipulation impairs performance only at long delays, but not at short delays, these findings would suggest that the impairment is likely due to a working memory deficit. Alternatively, a manipulation may impair both strategy selection and working memory. Because of the strategic component to the task, systematically examining the effects of different delays in a spatial delayed alternation or a delayed nonmatching-to-sample task can better elucidate whether a manipulation selectively impairs working memory.

To determine whether cholinergic transmission in the prefrontal cortex influences working memory, most studies have used intracranial drug infusions. Brito and colleagues (Brito, Silva, & Brito, 1989) demonstrated that injections of the muscarinic cholinergic antagonist scopolamine into the prelimbic area impaired performance in a T-maze, using a delayed nonmatching-to-sample procedure. Although the scopolamine infusions impaired performance, the effects of scopolamine were not examined across different delays, opening up the possibility that blockade of muscarinic receptors in the prelimbic area impaired the ability to use a delayed nonmatching-to-sample strategy, as opposed to a selective deficit in working memory. In a more recent experiment, blockade of muscarinic cholinergic transmission in the prelimbic area was found to impair both delayed nonmatching-to-sample and matching-to-sample in a spatial memory test using a 30-sec delay between the sample phase and the test phase (Granon et al., 1995). In this study, the effects of muscarinic cholinergic blockade were not examined at different delays either. However, the authors did report that pilot results indicated that prefrontal inactivation with the local anesthetic lidocaine did impair performance at a 30-sec delay, but not at a 5-sec delay (Granon et al., 1995). Because of the delay-dependent effects with lidocaine infusions, the impairment following scopolamine infusions into the prelimbic area may be due to a working memory deficit.

To better determine whether muscarinic cholinergic transmission in the prelimbic and infralimbic areas influences working memory processes, the effect of scopolamine infusions into the prelimbic–infralimbic areas across different delays was investigated recently in our laboratory with an experiment that used a spatial continuous recognition procedure (Ragozzino & Kesner, 1998). A continuous recognition procedure was developed in order to use a test that was comparable with the continuous recognition procedures employed to study working memory in humans. In this task, rats were exposed sequentially to 12 arms during a session. Three or 4 of the arms were presented for a second time in a session. A rat learned that a cereal reinforcement was at the end of the arm during the first presentation, but not during the second presentation. The number of intervening arm choices between the first and the second presentations of an arm varied from zero to six. Memory was assessed by the latency to enter an arm during the second presentation. Rats remember entering a location during a session and thus avoid entering the location during the second presentation. With an increase in lag, there is an increase in the delay and interference between the first and the second presentations of a spatial location; thus, latencies decrease with longer lags. Scopolamine injections produced a dose- and lag-dependent impairment in spatial working memory. More specifically, blockade of muscarinic receptors in the prelimbic–infralimbic areas did not impair memory at the shortest lag (Lag 0) but did impair performance with increased lags. Furthermore, the results also indicated that the deficit following scopolamine infusions increased at longer lags. The lag-dependent deficit is consistent with a working memory impairment, since scopolamine infusions into the prelimbic–infralimbic areas impaired spatial memory only at longer lags, when temporal and spatial interference was greatest. Moreover, the scopolamine-induced impairment was reversed with combined intracranial infusions of oxotremorine, a muscarinic agonist, suggesting that the impairment was specifically produced by blockade of muscarinic receptors in the prelimbic–infralimbic region. Overall, a continuous recognition procedure, such as that used in this study, has advantages in assessing memory performance across several different delays and levels of interference and may provide a useful paradigm for understanding the neurochemical mechanisms that underlie working memory.

In addition to spatial working memory tests in a T-maze or a radial arm maze, the effects of cholinergic drugs into the prelimbic area on matching-to-position tests in operant chambers have been investigated in other experiments (Aura & Riekkinen, 1999; Broersen, Heinsbroek, de Bruin, Uylings, & Olivier, 1995). In these tasks, rats were presented with one bar to press in a sample phase and then, after a delay, were presented with two bars in the test phase. Depending on the procedure in the test phase, a rat learned to press the bar that was the same as (matching-to-position) or the bar that was opposite to (nonmatching-to-position) that in the sample phase. Although this task has a spatial memory component, it is unclear to what degree the task involves allocentric spatial memory and/or egocentric spatial memory. More specifically, a rat may remember what bar it just pressed on the basis of extra-maze cues in the environment (allocentric), or a rat may remember on the basis of cues related to a particular body movement (egocentric). In accord with the findings in the radial arm maze, Broersen et al. (1995) demonstrated that scopolamine infusions into the prelimbic area pro-
duced a memory deficit in the matching-to-position task in a dose- and delay-dependent manner. In contrast, a recent study found that scopolamine injections into the prelimbic area did not impair delayed nonmatching-to-position performance (Aura & Reikkinen, 1999). However, in this experiment, only one dose of scopolamine was injected, which raises the possibility that the higher doses of scopolamine used in other studies that used a nonmatching-to-sample procedure and found memory deficits (Brito et al., 1989; Broersen et al., 1995; Granon et al., 1995) might have produced a working memory impairment.

The findings from experiments employing intracranial drug infusions on working memory tasks provide evidence that muscarinic cholinergic transmission in the prelimbic and infralimbic areas is important for working memory (Granon et al., 1995; Ragozzino & Kesner, 1998). This is further supported by results indicating that blockade of muscarinic receptors in the prelimbic area impairs working memory but does not impair visuospatial discrimination in a T-maze or a radial arm maze (Brito et al., 1989; Granon et al., 1995; Ragozzino & Kesner, 1998). For example, scopolamine infusions into the prelimbic–infralimbic areas do not impair performance when a rat must remember always to enter one spatial location for a reinforcement and to avoid entering another spatial location. Furthermore, acetylcholine release in the prelimbic area does not change from basal levels when rats perform a visual discrimination task (Himmelheber, Sarter, & Bruno, 1997). These findings suggest that cholinergic transmission—and, in particular, muscarinic cholinergic transmission—in the prelimbic–infralimbic areas is not critical for retention of a discrimination, as well as perceptual, motivational, or motor functions, but is important for spatial working memory.

Compared with muscarinic receptors, few studies have examined the role that nicotinic cholinergic receptors in the prelimbic–infralimbic areas play in working memory. One experiment in which the effects of nicotinic cholinergic transmission in the prelimbic area on working memory were investigated found that blockade of nicotinic receptors impairs spatial working memory in a delayed matching-to-sample procedure, but not in a delayed nonmatching-to-sample procedure (Granon et al., 1995). The reason for this selective deficit is unclear. Poucet and colleagues (Granon et al., 1995) have proposed that because a matching-to-sample rule requires a rat to go against its tendency to alternate, this procedure may be more difficult in that it makes greater attentional demands and/or requires a rat to shift strategies. Manipulations of the prefrontal area have been found to impair performance on tasks that either accentuate attentional processes or require a shift in strategies (de Bruin, Sánchez-Santed, Heinsbroek, Donker, & Postmes, 1994; Muir et al., 1996; Ragozzino, Detrick, & Kesner, 1999a; Ragozzino, Wilcox, et al., 1999). On the basis of the performance of control rats, there is a trend for the matching-to-sample procedure to be more difficult than the nonmatching-to-sample procedure (Granon et al., 1995). However, future experiments are needed in which changes in attentional demands and strategy selection, as well as their interactions, are systematically examined to better determine what role nicotinic cholinergic receptors in the prefrontal area play in cognitive processes.

**Cholinergic Influences in the Prelimbic and Infralimbic Areas on Attention**

Damage to basal forebrain cholinergic neurons impairs performance on different attention tasks, and because these neurons project to the medial prefrontal cortex, it has been assumed that the medial prefrontal cortex is also involved in some aspect of attention (Chiba, Bucci, Holland, & Gallagher, 1995; McGaughy, Kaiser, & Sarter, 1996; Muir, Everitt, & Robbins, 1994; Muir, Page, Sirinathsinghji, Robbins, & Everitt, 1993; Robbins et al., 1989; Stoehr et al., 1997; Turchi & Sarter, 1997). Indeed, medial prefrontal cortex lesions, centered in the prefrontal area, have been found to impair attention (Muir et al., 1996). The deficits following lesions centered in the prefrontal area were observed on a five-choice serial reaction time task that required sustained attention to a brief light stimulus across five different spatial locations. These lesions impaired choice accuracy and increased perseverative responding. Furthermore, in the lesioned rats, correct response latency increased, and there was a strong trend for accuracy to decline when attentional demands were augmented.

A series of recent experiments by Givens and colleagues (Gill, Masters, Sarter, & Givens, 1999; Gill, Sarter, & Givens, in press; Williams, Mohler, & Givens, 1999) examined the role of cholinergic input to different medial prefrontal cortex subregions in attentional processes. In one study, rats were trained to press one of three bars when a brief light stimulus appeared above it (Williams et al., 1999). The light stimulus occurred for a 200- or 500-msec duration and was presented in either a block of predictable trials, where the light stimulus was presented for the same duration for all the trials, or a block of random trials, where the light stimulus was presented for either 200 or 500 msec in a random fashion. Scopolamine infusions into the prefrontal area reduced choice accuracy in the random condition, but not in the predictable condition, suggesting that muscarinic cholinergic transmission in the prefrontal area is important under conditions in which there are increased attentional demands (Williams et al., 1999). These results are comparable with the lesion study discussed earlier, which demonstrated that lesions centered in the prefrontal area impaired response selection with enhanced attentional demands (Muir et al., 1996).

In another study by this group, recordings of prefrontal single-unit activity during sustained attention performance before and after unilateral cholinergic deafferrntation of the medial prefrontal cortex were examined (Gill et al., in press). In this experiment, rats were tested on a visual stimulus detection task. If a visual stimulus (25, 50, or 500 msec in duration) was presented on a trial, a rat had to press one of two bars in an operant chamber to re-
receive a water reinforcement. If no visual stimulus was presented on a trial, a rat had to press the other bar. During certain trials, an overhead flashing light was presented as a visual distractor to increase attentional demands. Several neurons in the prelimbic region exhibited correlated activity either when the rats were preparing to make a response or during response emission, consistent with the idea that the prefrontal cortex is involved in the planning and execution of goal-directed behavior. In addition, an even greater number of prelimbic neurons showed correlated activity in anticipation of a reward and during consumption of a reward (Gill et al., in press). In contrast, unit activity did not correlate with the presentation of sensory stimuli. Nevertheless, presentation of the visual distractor did lead to a decrement in performance and a simultaneous altering of the firing rate of several prefrontal cells.

Injecting the cholinergic neurotoxin 192 IgG-saporin into the medial prefrontal cortex produced a significant reduction in the response- and reward-correlated activity of these prelimbic neurons. Cholinergic deafferentation also decreased the proportion of neurons modulated by the presentation of the distractor. Moreover, those neurons that did change their activity in the presence of the distractor showed a reduced magnitude of change, as compared with those recorded prior to the cholinergic lesion. However, although a unilateral cholinergic lesion of the medial prefrontal cortex significantly altered the response properties of prelimbic neurons during a sustained attention task, this lesion did not impair task performance. More recently, preliminary results from a subsequent study indicate that bilateral infusions of 192 IgG-saporin into the medial prefrontal cortex do impair sustained attention (Gill et al., 1999). Taken together, these results suggest that neuronal activity in the prelimbic area is related to sustained attention and that cholinergic modulation of cellular firing in this subregion may be important for accurate performance on tasks requiring sustained attention (Gill et al., 1999; Gill et al., in press). Important to note, however, because the neurotoxin was infused into multiple subregions within the medial prefrontal cortex, areas outside the prelimbic area may also have contributed to the impairing effects of cholinergic deafferentation on attentional processing.

Nonetheless, experiments measuring acetylcholine output by in vivo microdialysis with high pressure liquid chromatography also implicate prelimbic cholinergic afferents in attention. In particular, a study in which acetylcholine output was assessed in the prelimbic area during a sustained attention task found that acetylcholine output increased when the attentional demands were increased by adding a flashing light during the task (Sarter et al., 1996). This result contrasts with findings discussed earlier that acetylcholine output in the prelimbic area did not change when a flashing light was presented during a “simple” discrimination task (Himmelheber et al., 1997). In the sustained attention task, a rat had to differentially respond to receive a reinforcement, depending on the type of signal presented. However, in the discrimination task, the rule was to execute the same response when a signal appeared. Thus, one possibility is that the differential response selection task required greater attentional processing and, concomitantly, higher levels of acetylcholine in the prelimbic area (Sarter et al., 1996).

Changes in acetylcholine output were also observed in other conditions not directly related to task performance (Sarter et al., 1996). Acetylcholine output was enhanced when rats were moved into the test chamber and at the initial onset of a test session (Sarter et al., 1996). This increase in acetylcholine release may be related to attention or arousal mechanisms activated when animals are moved to a different environment.

Cholinergic Influences in the Anterior Cingulate on Working Memory

In one recent experiment, the effects of cholinergic drug infusions into the anterior cingulate on working memory were assessed in a spatial memory task using a 12-arm radial maze. In contrast to infusions into the prelimbic areas, injections of scopolamine into the dorsal anterior cingulate did not impair working memory in a spatial continuous recognition test (Ragozzino & Kesner, 1998). In another experiment, the effects of muscarinic cholinergic receptor blockade in the dorsal anterior cingulate during a delayed matching-to-position task was determined. In this task, rats were tested in an operant chamber that had two retractable levers. During the sample phase of a trial, one of the two levers was presented. After a response on the sample lever, the level was retracted. Following a delay, both levers were presented during the test phase. To receive a reinforcement, a rat had to press the same lever as that in the sample phase. Scopolamine injections into the dorsal anterior cingulate impaired performance on a delayed matching-to-position task, but the deficit was delay independent, suggesting that the muscarinic receptors in this area were not specifically critical for working memory performance (Herremans, Hjizen, & Olivier, 1997; Herremans, Welborn, Hjizen, Olivier, & Slangen, 1996). Similarly, Pepeu and colleagues (1996) measured acetylcholine output, using a transverse dialysis probe centered in the dorsal and ventral anterior cingulate. It was demonstrated that acetylcholine release was increased during spontaneous alternation performance in a Y-maze, a task that has a spatial working memory component (Ragozzino, Unick, & Gold, 1996). However, it is unclear whether the change in acetylcholine output was related to working memory, novelty, or some combination. Acetylcholine levels were measured in this study during the rat’s first exposure to the maze, and an increase in acetylcholine release is commonly observed in several forebrain regions when rats are placed in a novel or different environment (Acquas, Wilson, & Fibiger, 1996; Aloisi, Casamenti, Scalì, Pepeu, & Carli, 1997; Orsatti, Casamenti, & Pepeu, 1996; Sarter et al., 1996).

When evaluated in conjunction with other studies indicating that anterior cingulate lesions do not impair spatial working memory (Ragozzino et al., 1998; Sánchez-Santed, de Bruin, Heinsbroek, & Verwer, 1997; Silva...
et al., 1986), these results suggest that cholinergic trans-
mission in the anterior cingulate does not modulate spatial
working memory processes, as was observed in the pre-
limbic and infralimbic areas (Herremans et al., 1997; Her-
remans et al., 1996; Ragozzino & Kesner, 1998). Rather,
the findings raise the possibility that cholinergic input to
the medial prefrontal cortex modulates different cognitive
processes determined by the particular prefrontal cortex
subregion. Thus, different prefrontal cortex subregions
may differentially contribute to various cognitive functions
that can be modulated by cholinergic input to those sub-
regions.

Cholinergic Influences in the
Anterior Cingulate on Attention

The behavioral effects of muscarinic cholinergic block-
ade in the anterior cingulate on different discrimination
tasks appear to be more related to changes in attention,
as opposed to working memory (Herremans et al., 1997;
Herremans et al., 1996). More specifically, Herremans
and colleagues found that scopolamine infusions into the
anterior cingulate during a matching-to-position or a
tone–light conditional discrimination task decreased dis-
criminability at no delay and appeared to produce a gen-
eral increase in distractability. In another experiment in
which the effects of scopolamine infusions into the ante-
cior cingulate on attention were investigated, it was found
that muscarinic cholinergic blockade in this subregion
impaired performance during a varying stimulus duration
condition, but not during a predictable stimulus duration
condition (Williams et al., 1999). Like the findings in the
prelimbic area, these results suggest that muscarinic
cholinergic transmission in the anterior cingulate is im-
portant when there is an increase in attentional demands
(Williams et al., 1999).

Further support for the idea that cholinergic input to the
anterior cingulate modulates attentional processes comes
from an experiment by Pirch and colleagues (Pirch, Turco,
& Rucker, 1992) in which single neurons in the anterior
cingulate were recorded during an auditory discrimina-
tion task. Neurons in the anterior cingulate were found to
exhibit discrimination-related responses to conditioned
tones, which were modulated by direct application of
acetylcholine. These acetylcholine-induced changes were
blocked by the muscarinic antagonist atropine (Pirch
et al., 1992).

In other experiments, acetylcholine output increased in
the anterior cingulate during exposure to novel sensory
stimuli or environments (Acquas et al., 1996; Giovannini,
Bartolini, Kopf, & Pepeu, 1998; Inglis & Fibiger, 1995).
Conversely, there was no change in acetylcholine output
after extensive exposure to unconditioned stimuli (Acquas
et al., 1996). Novel conditions are commonly interpreted
as highly arousing and may engage attentional processes
for exploring particular environments or conditions. Thus,
the changes in acetylcholine output in the anterior cin-
gulate during novel situations may reflect an increase in
attentional demands. Whether the novelty-induced changes
in acetylcholine release in different medial prefrontal
subregions are related to the same processes (i.e., atten-
tion) or to different processes is still to be determined.

THE ROLE OF DOPAMINERGIC
TRANSMISSION IN THE PREFRONTAL
CORTEX ON COGNITIVE PROCESSES

A few behavioral experiments have studied the influ-
ence of dopamine transmission in the medial prefrontal
cortex on working memory and learning. In addition to
traditional pharmacological approaches, in vivo micro-
dialysis, in vivo voltammetry, and the use of the cate-
cholamine neurotoxin 6-OHDA have provided approaches
for understanding how dopaminergic transmission in sep-
parate prefrontal cortex subregions influences different be-
havioral and cognitive processes.

Dopaminergic Influences in the Prelimbic
and Infralimbic Areas on Working Memory

Current findings suggest that dopamine afferents
in the prelimbic–infralimbic areas modulate working
memory. One of the first experiments to suggest a role of
prelimbic–infralimbic dopamine in working memory
demonstrated that dopamine depletion in the prelimbic–
infralimbic areas impaired delayed alternation in both a
T-maze and a radial arm maze (Busber & Schmidt, 1990).
In this experiment, the lesion-induced deficit was selec-
tive to working memory, since these same lesions did not
affect performance on alternation tests without delays. In
another study, Sokolowski and Salamone (1994) demon-
strated that dopamine depletion, by use of 6-OHDA, in the
prelimbic–infralimbic areas impaired performance on a
differential reinforcement of low rates of responding task
with a 30-sec schedule. In this task, a rat must withhold
responding until 30 sec elapses to receive a reinforce-
ment. This test involves memory for temporal duration and
likely recruits working memory processes. Although this
task has also been used to measure impulsivity, it is im-
portant to note in this study that lesioned rats were more
likely to respond before 30 sec had elapsed but did not
exhibit any motor abnormalities (i.e., hyperactivity),
suggesting that this deficit was selective to temporal dis-
crimination (Sokolowski & Salamone, 1994).

The results from investigations using intracranial drug
injections suggest that D1 receptors in these areas are
particularly critical for working memory performance.
In examining working memory with a delayed matching-
to-position procedure, infusions of SCH-23390, a dopa-
mine D1 antagonist, into the prelimbic area impaired per-
f ormance (Broersen et al., 1995). The drug-induced deficit
was dose dependent, and there was a trend for the impair-
ment to be delay dependent (Broersen et al., 1995). Block-
ade of D1 receptors in the prelimbic area has also been
found to produce spatial working memory deficits in a
dose-dependent manner (Seamans, Floresco, & Phillips,
1998). Infusions of a full D1 receptor agonist into the pre-
limbic area likewise produced spatial working memory
impairments (Zahrt, Taylor, Mathew, & Arnsten, 1997).
These results suggest that there is an optimal window of
dopamine D_1 receptor stimulation in the prelimbic area for facilitation of working memory and that either a paucity or an excess of prelimbic D_1 receptor stimulation may impair performance.

In contrast to D_2 receptors, infusions of the D_2 antagonist sulpiride into the prelimbic area do not impair working memory (Brito et al., 1989; Seamans et al., 1998). Unknown is what effect selective D_3, D_4, and D_5 agents infused into the prelimbic–infralimbic areas would have on working memory. Thus, to date it appears that dopamine stimulation of prelimbic D_1 receptors plays a role in working memory and that there is a limited range of D_1 receptor stimulation that is optimal for working memory performance.

**Dopaminergic Influences in the Prelimbic–Infralimbic Areas on Learning**

A recent experiment examined the effects of 6-OHDA lesions centered in the prelimbic–infralimbic areas on acquisition and expression of conditioned fear, using tone–shock pairings (Morrow, Elsworth, Rasmusson, & Roth, 1999). Dopamine depletion did not impair the learning or expression of a conditioned fear response. However, dopamine depletion did impair the extinction of fear conditioning. These findings suggest that dopamine input to the prelimbic–infralimbic areas may be critical for behavioral flexibility—that is, adapting behavior to changes in environmental conditions, rather than the learning of conditioned fear. Further support for this idea comes from a preliminary study in which the effects of D_1 receptor blockade in the learning and reversal of behavioral rules were examined (Ragozzino, Detrick, & Kesner, 1999b). Blockade of D_1 receptors in the prelimbic–infralimbic areas did not impair learning of a visual cue or response (turn) discrimination in a cross-maze but did impair performance when rats had to shift from one discrimination strategy to the other discrimination strategy (Ragozzino, Detrick, & Kesner, 1999b).

In contrast to the experiments described above, the findings from in vivo microdialysis studies during acquisition suggest a role for prelimbic–infralimbic dopamine release in learning. In assessing acquisition of conditioned fear, dopamine was observed to increase during the presentation of a stimulus (tone or flashing light) not paired with footshock, with further increases in dopamine release when the stimulus was paired with footshock (Wilkinson et al., 1998). This change in dopamine output diminished across conditioning trials in conjunction with enhanced freezing, which was used as a measure of conditioned fear. These findings suggest that dopamine release in the prelimbic–infralimbic areas contributes to new learning.

Dopamine output was also found to increase during acquisition of a barpress operant under a continuous reinforcement schedule (Izaki, Hori, & Nomura, 1998). The observed increase in dopamine release across learning contrasts with the experiment by Wilkinson et al. (1998), which found the opposite pattern during conditioned fear acquisition. Because the tasks had different types of reinforcement (aversive vs. appetitive), one possibility is that the dynamics of dopamine release for learning an aversive task is different from that in an appetitively motivated task. During a retention session in the operant task, there was no change in dopamine output from basal levels (Izaki et al., 1998). These results support the idea that dopamine output in the prelimbic–infralimbic areas is related to stimulus–reward learning, but not for retrieval of the learned association.

Although the results from studies suggesting that prelimbic–infralimbic dopamine release is involved in learning appear to conflict with experiments demonstrating that 6-OHDA lesions or dopamine receptor blockade do not impair acquisition, the two sets of findings may actually be related. More specifically, in the conditioned fear study in which dopamine output was concomitantly measured with freezing behavior, rats were exposed to the test apparatus for several minutes without any exposure to shock (Wilkinson et al., 1998). Subsequently, rats were presented a stimulus with footshock. The increase in dopamine release with the first presentation of the stimulus–shock pairing may be related to the elicitation of a new response pattern induced by changes in the environmental conditions. Similarly, in the operant task, rats were exposed to the operant chamber on several sessions, receiving a reinforcement under a variable interval schedule without barpress training (Izaki et al., 1998). During the acquisition session, rats now had to press a bar to receive a reinforcement under a continuous reinforcement schedule. Thus, in this experiment, the conditions were also changed, requiring a different response pattern. Taken together, the findings suggest that changes in dopamine release in the prelimbic–infralimbic areas may underlie the flexible shifting of a learning set to changes in environmental demands.

The idea that dopamine output plays a role in behavioral flexibility is further supported by a recent study using in vivo voltammetry (Richardson & Gratton, 1998). In vivo voltammetry has advantages over in vivo microdialysis in that dopamine changes can be observed in the millisecond range versus the minute range, and the spatial resolution is also better. In this study, rats were trained to barpress for a liquid reinforcement under different reinforcement schedules. Changes in dopamine release in the prelimbic or the infralimbic area occurred when the temporal component of the learned association was changed or the reinforcement that a rat expected was changed. Dopamine release did not change during presentation of the conditioned stimuli or consumption of the liquid reinforcement. Thus, dopamine release selectively changed when the learning conditions were modified (Richardson & Gratton, 1998). These findings provide additional evidence that dynamic changes in prelimbic dopamine release may underlie behavioral flexibility.

**Dopaminergic Influences in the Anterior Cingulate on Working Memory**

In one experiment, the effects of SCH-23390 infusions into the anterior cingulate on a delayed matching-
to-position task in an operant chamber were examined. The procedure was the same as that described earlier. Blockade of D1 receptors did not impair working memory performance (Herremans et al., 1996). Importantly, the drug doses used in this experiment were doses previously found to affect behavioral performance when injected into other brain sites, suggesting that activation of dopaminergic receptors in the anterior cingulate does not influence working memory. Moreover, these findings are consistent with lesion studies demonstrating that anterior cingulate lesions do not impair working memory (Ragozzino et al., 1998; Sánchez-Santed et al., 1997; Silva et al., 1986).

**SUMMARY AND CONCLUSIONS**

In recent years, several experiments have attempted to unravel how cholinergic and dopaminergic afferents in different subregions of the rat prefrontal cortex contribute to learning, memory, and attention. Activation of cholinergic or dopaminergic receptors in the prelimbic and infralimbic areas facilitates working memory. In particular, numerous findings suggest that activation of muscarinic cholinergic or activation of D1 dopamine receptors in the prelimbic–infralimbic areas contributes to working memory. In contrast, stimulation of D2 receptors in the prelimbic–infralimbic areas does not appear to play a significant role in working memory. Less is known about the contribution of nicotinic cholinergic receptors to working memory.

More extensive research is needed to determine whether acetylcholine and dopamine directly interact within the prefrontal cortex to facilitate working memory or whether their actions are independent. Some experiments suggest an interaction between acetylcholine and dopamine in the medial prefrontal cortex, whereas other studies do not (Moore, Fadel, Sarter, & Bruno, 1999; Yang & Mogenson, 1990).

Cholinergic input to the prelimbic–infralimbic areas may also influence attentional processing, particularly when there is an increase in attentional demands. However, it is possible that cholinergic input more generally influences behavioral flexibility, only incidently affecting attentional processes. This is based on previous findings indicating that medial prefrontal lesions, centered in the prelimbic–infralimbic areas, produced random barpressing, an effect not influenced by manipulations that increase attentional demands (Miner, Ostrander, & Sarter, 1997).

Activation of dopamine receptors in the prelimbic–infralimbic areas may also contribute to behavioral flexibility. Different lines of evidence suggest that there are dynamic changes in dopamine release in the prelimbic–infralimbic areas when there are changes in environmental demands. In addition, preliminary evidence suggests that activation of D1 receptors may facilitate behavioral flexibility or strategy switching.

In contrast to the prelimbic and infralimbic areas, the cholinergic and dopaminergic input to the anterior cingulate does not appear to play a critical role in spatial working memory. Rather, cholinergic input to the anterior cingulate seems intimately involved in attentional processing. More specifically, activation of muscarinic cholinergic receptors occurs when there is an increase in attentional demands. These results also suggest that whatever attentional requirements there are in most working memory tests, they are different from the attentional processes mediated by the anterior cingulate. Examining the role of nicotinic and dopamine receptors will be important to building a more complete picture of cholinergic and dopaminergic modulation of anterior cingulate activity related to attention.

Overall, the findings suggest that acetylcholine and dopamine modulate activity in the medial prefrontal cortex to facilitate different behavioral and cognitive processes. Importantly, current findings, particularly in the prelimbic and infralimbic regions, suggest that the same prefrontal cortex subregions may contribute to various functions—that is, working memory and strategy selection—under different dynamic states and that cholinergic and dopaminergic afferents influence these different states of activity. However, there is still the possibility that the diverse behavioral deficits produced by manipulations of the prelimbic–infralimbic areas represent disruption of a single process (i.e., attention). Future experiments will be important in further elucidating the role of prefrontal cholinergic and dopaminergic afferents in modulating cognitive processes.

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