Differential effects of intra-amygdala lidocaine infusion on memory consolidation and expression of a food conditioned place preference

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In the present experiments, we examined the effects of posttraining and preretention intra-amygdala infusion of the anesthetic lidocaine on memory consolidation and expression in a food conditioned place preference (CPP) task. In two separate experiments, for 4 alternating days, food deprived adult male Long-Evans rats were given access to food or no access to food during confinement for 30 min to one of two compartments in a place preference apparatus. On Day 9, the rats were given a 20-min test session and allowed access to all compartments of the apparatus. No food was present on the test day, and the amount of time spent in each compartment of the apparatus was recorded. On the test day, rats receiving saline immediately after previous training, or immediately prior to testing, spent more time in the compartment that was previously paired with food, demonstrating a food conditioned place preference (CPP). Posttraining infusion of lidocaine (2% solution, 0.5 μl/side) into the basolateral amygdala blocked acquisition, indicating that this structure is necessary for memory consolidation processes that mediate a food CPP. In contrast, immediate pretest infusion of lidocaine into the basolateral amygdala had no effect on expression of a food CPP. Taken together, these findings suggest a modulatory role for the basolateral amygdala in memory consolidation processes that underlie reward-related learning in a food CPP task.

The conditioned place preference (CPP) paradigm is commonly used to infer the rewarding properties of various substances by assessing the extent to which stimuli that have been paired with treatment administration elicit approach responses in the absence of the treatment on the test day (for reviews, see Carr, Fibiger, & Phillips, 1989; Schechter & Calcagnetti, 1993). Drugs of abuse including amphetamine, cocaine, diazepam, nicotine, and morphine, as well as naturally occurring stimuli such as food, water, or exposure of male rats to a receptive female, produce CPPs (Carr et al., 1989; Schechter & Calcagnetti, 1993).

One overlooked aspect of the CPP paradigm is that it offers an opportunity to assess not only the development of primary reward, but also memory for the rewarding consequences of various treatments (White & Carr, 1985). Specifically, because subjects are in a treatment-free state on the test day, expression of a CPP requires memory for the association between environmental cues and the affective state produced by the treatment. An important role of the memory enhancing properties of treatments in the CPP paradigm was demonstrated in a study examining the rewarding effects of glucose and saccharin. Both of these substances are orally consumed in preference tests, suggesting that on the basis of their gustatory input, they both possess rewarding affective properties (White & Carr, 1985). However, glucose, but not saccharin, has the additional properties of producing a CPP following systemic injection and enhancing memory in several tasks when administered posttraining (e.g., Gold, 1986; Messier & White, 1984; Packard & White, 1990; White & Carr, 1985). Importantly, although systemic injections of saccharin are not capable of producing a CPP, additional posttraining administration of glucose has been shown to induce a saccharin CPP (White & Carr, 1985). Taken together, these findings suggest that acquisition of a CPP is dependent on both a treatment’s ability to produce a rewarding affective state and its memory enhancing effect. Consistent with this hypothesis, other substances that produce a CPP (e.g., amphetamine and cocaine), also enhance memory when injected posttraining (e.g., Cestari, Mele, Oliverio, & Castellano, 1996; Doty & Doty, 1966; Packard & White, 1989). In addition to memory enhancement, postraining treatments can be used to block or attenuate memory in the CPP paradigm. For example, intracerebroventricular infusion of protein kinase inhibitors block a cocaine CPP (Cervo, Mukherjee, Bertaglia, & Samanin, 1997).

Recent studies examining the neuroanatomical bases of reward-related learning and memory as measured in the CPP task have implicated the basolateral/lateral amygdala. For example, pretraining quinolinic acid lesions of the amygdala block acquisition of a cocaine CPP (Brown & Fibiger, 1993), and electrolytic and excitotoxic lesions of the lateral amygdaloid nucleus prevent the acquisition and expression of an amphetamine CPP (Hiroi & White, 1986).
1991). In addition to psychostimulant drug treatments, the use of naturally occurring stimuli in the CPP task has also implicated amygdala function in CPP behavior. Thus, pretraining electrolytic lesions of the lateral nucleus of the amygdala block acquisition of a food CPP in the eight-arm radial maze (McDonald & White, 1993), and acquisition/expression of a CPP for sucrose is abolished by excitotoxic lesions of the basolateral amygdala (Everitt, Morris, O'Brien, & Robbins, 1991). Finally, pretraining infusion of the acetylcholine antagonist scopolamine into the amygdala prevents acquisition of a food CPP (McIntyre, Ragozzino, & Gold, 1998).

Although previous findings clearly implicate the amygdala in CPP behavior, the extent to which these manipulations influence memory processes in the CPP task can be difficult to assess in studies that employ pretraining or pretest lesion techniques. Such treatments may influence CPP acquisition and/or expression by impairing memory, primary reward, or nonmemonic sensory, motivational, or motoric processes. Although the contribution of nonmemonic factors in amygdala mediation of reward-related learning have been indirectly addressed in studies that have revealed dissociations following amygdala lesions in different types of learning tasks (e.g., Burns, Everitt, & Robbins, 1993; Everitt et al., 1991; McDonald & White, 1993), the present experiments were designed to directly examine the role of this structure in memory processes underlying a food CPP. Specifically, saline or the local anesthetic lidocaine were infused immediately posttraining into the basolateral amygdala to examine the role of this structure in memory consolidation processes (McGaugh, 1966, 1973, 1989) mediating a food CPP.

In addition to examining the effects of posttraining intrabasolateral amygdala infusion of lidocaine, the question of whether the amygdala plays a differential role in memory consolidation and expression of a food CPP was assessed. This question is of interest in view of evidence that, for at least some learning tasks, posttraining amygdala activation modulates memory storage processes that occur in other brain sites (e.g., hippocampus and caudate-putamen), rather than as its serving as a site of permanent memory storage (Packard, Cahill, & McGaugh, 1994; Packard, Cahill, Williams, & McGaugh, 1995; Packard & Teather, 1998). Evidence from studies employing permanent lesion techniques indicates a role for the amygdala in CPP expression for both amphetamine (lateral amygdala; Hiroi & White, 1991) and sucrose (basolateral/lateral amygdala; Everitt et al., 1991). In an additional experiment, saline or lidocaine was infused into the basolateral amygdala prior to the test session to determine whether this structure also mediates expression of a food CPP.

METHOD

Subjects

The subjects were 30 adult male Long-Evans rats (300–375 g). The rats were individually housed in a temperature-controlled environment on a 12:12-h light:dark cycle with the lights on from 7 am to 7 pm and were given ad-lib access to food and water prior to surgery.

Apparatus

The apparatus was constructed of wood and had a Plexiglas front wall. It consisted of three compartments, two of which were identical in size (45 x 45 x 30 cm high). One compartment was painted black and had a black Plexiglas floor and 1 ml of a 1% acetic acid solution sprayed on the walls prior to training and testing. The other compartment was painted white and had a white Plexiglas floor. These two compartments were separated from each other by a wooden partition and were connected by the third compartment. The gray roofed tunnel (36 x 18 x 20 cm high) had a metal floor and could be opened to allow entrance into each of the two larger compartments.

Surgery

Prior to surgery, the subjects were anesthetized with a cocktail of 30-mg/kg ketamine HCl and 2.5-mg/kg xylazine. Bilateral guide canulae (15 mm in length) were implanted overlying the basolateral amygdala by 1 mm, using standard stereotaxic techniques. Jeweler's screws were anchored to the skull and attached to the cannulae with dental acrylic. Stereotaxic coordinates for the basolateral amygdala placements were AP = -2.2 mm, ML = 4.7 from bregma, and DV = -7.0 mm from the skull surface. After surgery, stylets were inserted and left in the cannulae until the time of injection. The subjects were allowed 1 week of recovery before food deprivation was initiated.

Injection Procedures

The intra-amygdala infusions were administered via a microsyringe pump using a 10-μl Hamilton syringe connected to polyethylene tubing. The infusions were administered over a 53-sec period, and the injection needles were left in place an additional 60 sec to allow for diffusion. Injection volume was 0.5 μl for both saline and lidocaine (2% solution, 10 μg/side) injections. This dose and volume of lidocaine was chosen on the basis of our previous findings, which indicated that it is behaviorally effective when administered into the amygdala (Packard & Chen, 1999; Packard & Teather, 1998). Electrophysiological evidence indicates that intracerebral injection of this concentration of lidocaine solution depresses neural function within 2 min of infusion, with a duration of approximately 30 min (Tehovnik & Sommer, 1997). This consideration is important in view of our use of a 20-min session on the test day in the CPP task.

Histology

Following behavioral testing, the animals were deeply anesthetized with a 1-ml injection of sodium pentobarbital (50 mg/kg) and perfused with saline followed by 10% formol-saline. The brains were removed and sectioned at 20 μm. They were then stained with cresyl violet, and cannula placements were verified using the atlas of Paxinos and Watson (1986). As illustrated in Figure 1, injection needle tips were located in the basolateral nucleus of the amygdala, ranging from -2.12 to -2.8 mm AP from bregma.

Behavioral Procedures

One week following surgery, the rats were reduced to 85% of their ad-lib feeding weight over an additional 7-day period. CPP training took place over 6 days, and the parameters were chosen on the basis of pilot data indicating that they produced a significant food CPP. The 1st day consisted of habituation, during which the subjects were allowed access to all three compartments of the CPP apparatus for 10 min. The next 4 days consisted of two food and two nonfood pairings, during which the subjects were confined to a compartment containing 7 g of Noyes food pellets scattered on the floor, or no food. Half of the subjects had access to food during exposure.
Figure 1. Location of cannula placements in the basolateral amygdala. The placements ranged from $-2.12$ to $-2.8$ mm anterior-posterior to bregma (Paxinos & Watson, 1986) for both infusions made posttraining (left column) and pretesting (right column).

to the black compartment and no access to food during exposure to the white compartment. The remaining subjects had access to food during exposure to the white compartment and no access to food during exposure to the black compartment. Half of the subjects had access to food on odd numbered days, and half had access to food on even numbered days. Following the 4 training days, the subjects were given a 20-min test session during which they were allowed access to all three compartments of the CPP apparatus with no food present. Time spent in each of the compartments was recorded.

To examine the effects of posttraining intrabasolateral amygdala infusion of lidocaine, the subjects received bilateral infusions of lidocaine ($n = 6$) or saline ($n = 8$) immediately following removal from their paired (i.e., food) and unpaired compartments on each of the training days, and on the test day, the subjects were allowed access to all three compartments of the apparatus with no food present. To examine the effects of pretesting infusions of lidocaine, the subjects received bilateral infusions of lidocaine ($n = 8$) or saline ($n = 8$) immediately prior to the 20-min test session in the food CPP paradigm.

Results

Posttraining intra-amygdala lidocaine infusions. Figure 2 shows the amount of time spent in the paired and unpaired compartments on the test day. A two-way one repeated measure analysis of variance (ANOVA) with posttraining infusion as an independent variable and paired versus unpaired side as a repeated measure revealed a significant interaction $[F(1,12) = 26.48, p < .01]$. Post hoc two-tailed paired $t$ tests revealed that the subjects receiving posttraining intrabasolateral amygdala saline infusions spent significantly more time in the food paired compartment than in the nonfood paired compartment $[t(7) = 6.30, p < .05]$, indicating a food CPP. In contrast, the rats receiving posttraining intrabasolateral amygdala lidocaine infusions did not spend different amounts of time in the food and nonfood paired compartments $[t(5) = 1.27, n.s.]$. These findings indicate a blockade of a food-CPP by posttraining infusions of lidocaine into the basolateral amygdala.

Pre-test intra-amygdala lidocaine infusions. Figure 3 shows the amount of time spent in the paired and unpaired compartments on the test day. A two-way one repeated measure ANOVA with posttraining infusion as
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Figure 2. Effect of posttraining intrabasolateral amygdala infusion of lidocaine on place-memory consolidation of a food conditioned place preference. Mean time spent (second ±SEM) during a 20-min test session in the previously paired (food) and unpaired (no food) compartments of the place preference apparatus. *Significant place preference.

an independent variable and paired versus unpaired side as a repeated measure revealed no significant interaction \( F(1,14) = 0.45, \text{n.s.} \). A significant effect of side \( F(1,14) = 21.1, p < .01 \) indicated a significant food CPP in rats infused with either saline or lidocaine immediately prior to testing.

DISCUSSION

The findings indicate that posttraining infusions of lidocaine into the basolateral amygdala block memory consolidation processes mediating a food CPP. This finding is consistent with the impairing effects of pretraining basolateral/lateral amygdala lesions on acquisition of drug (Brown & Fibiger, 1993; Hiroi & White, 1991), sucrose (Everitt et al., 1991), and food-induced (McDonald & White, 1993) CPPs. However, in previous studies using pretraining amygdala lesions, the impairing effects observed in CPP behavior may have been due to an influence on learning and/or memory per se, or on nonmnemonic factors such as primary reward, sensory processes, or motivation. The contribution of such factors to amygdala lesion-induced impairments in CPP behavior appear unlikely in view of evidence that similar lesions do not effect acquisition of other types of learning tasks with similar nonmnemonic characteristics (e.g., Cador, Robbins, & Everitt, 1989; Everitt et al., 1991; McDonald & White, 1993). Nonetheless, our findings using a reversible posttraining lesion technique clearly suggest a role for the basolateral amygdala in memory processes underlying a food CPP (i.e., consolidation; McGaugh, 1966, 1973, 1989). Moreover, they raise the possibility that the impairment produced by pretraining lesions are due, at least in part, to the absence of a functional amygdala during the posttraining period.

Pretest infusion of lidocaine into the basolateral amygdala did not block expression of a food CPP, suggesting that in this task, expression of approach responses to conditioned stimuli associated with the rewarding affective properties of food does not require basolateral amygdala function. Electrophysiological evidence indicates that intracerebral injection of a 2% lidocaine solution depresses neural function within 2 min of infusion and has a duration of approximately 30 min (Tehovnik & Sommer, 1997), suggesting that neural inactivation was present during the 20-min test session in our CPP task. However, our findings stand in contrast to those of previous studies examining the effects of excitotoxic or electrolytic lesions of the basolateral/lateral amygdala on expression of various CPPs. For example, pretest quinolinic acid lesions of the basolateral amygdala prevented expression of a sucrose CPP (Everitt et al., 1991), and electrolytic or NMDA-induced lesions of the lateral amygdala prior to the test day blocked expression of an amphetamine CPP (Hiroi & White, 1991). Although the use of excitotoxic lesions have the important advantage of sparing neuronal fibers of passage, the amygdaloid complex is highly susceptible to seizure activity, and excitotoxic lesions of the amygdala have been found to cause damage to associated limbic and cortical structures (e.g., Berger, Charton, & Ben-Ari, 1986). This raises the possibility that extra-amygdala damage (e.g., perirhinal cortex, as described in the study of Everitt et al., 1991) may have contributed to the behavioral impairments observed during CPP expression. In the latter study (Hiroi & White, 1991), blockade of expression of an amphetamine CPP was observed with lesions primarily restricted to the lateral amygdala. It is possible that expression of an amphetamine, but not a food CPP requires amygdala function. Alternatively, lidocaine infusion into the basolateral site targeted in the present study may not have adequately inactivated more lateral regions of the amygdala that could conceivably be involved in the expression of a

Figure 3. Effect of pretesting intrabasolateral amygdala infusion of lidocaine on expression of place memory of a food conditioned place preference. Mean time spent (second ±SEM) during a 20-min test session in the previously paired (food) and unpaired (no food) compartments of the place preference apparatus. *Significant place preference.
food CPP. According to this view, given that our basolateral injection site was effective when lidocaine was infused posttraining, it is possible that the basolateral amygdala modulates the lateral amygdala during CPP consolidation, and that the lateral amygdala is therefore necessary for CPP expression. However, although our injection sites were within the basolateral nucleus, in view of the injection volume used (0.5 µl), it is unlikely that lidocaine selectively affected this amygdaloid nucleus. Further studies targeting different amygdala nuclei and employing smaller injection volumes are clearly necessary to examine the possibility of a differential role of amygdaloid nuclei in mediating memory consolidation and expression of CPP behavior.

Taken together, the present patterns of results, in which intrabasolateral infusions of lidocaine block memory consolidation processes but not behavioral expression of a food CPP, suggest either multiple storage sites (i.e., amygdala and other brain sites) for stimulus reward associations, or a temporary nonstorage “modulatory” role for the amygdala in memory for conditioned reward. These findings are similar to those of studies demonstrating a time-dependent role of the amygdala in memory in other types of learning tasks (e.g., McGaugh, Cahill, & Roozendaal, 1996; Packard et al., 1995; Parent, West, & McGaugh, 1994; Poremba & Gabriel, 1999). For example, posttraining intra-amygdala injections of amphetamine enhance memory in hippocampal-dependent and caudate nucleus-dependent hidden and visible platform water maze tasks (Packard et al., 1994; Packard & Teather, 1998), respectively. Importantly, the memory enhancement produced by intra-amygdala amphetamine in these tasks is not due to storage of memory within this structure, since pretreatment intra-amygdala injections of lidocaine do not prevent expression of amphetamine’s memory enhancing effects (Packard et al., 1994; Packard & Teather, 1998). In contrast, concurrent posttraining infusion of lidocaine into the hippocampus selectively blocks the memory enhancing effects of intra-amygdala injection of amphetamine in a hidden platform task, whereas similar infusion of lidocaine into the caudate-putamen selectively blocks the memory enhancing effects of intra-amygdala injection of amphetamine in a visible platform task (Packard & Teather, 1998). Thus, independent hippocampal and caudate-putamen memory systems, which both receive efferent basolateral amygdala projections (Finch et al., 1986; Kita & Kitai, 1990; Krettek & Price, 1977, 1978; McDonald, 1991; Russchen & Price, 1984), appear to be two targets of the memory modulatory influence of the amygdala.

An important question concerns identification of the brain region(s) that may receive a memory modulatory influence from the basolateral amygdala in the CPP task. Previous evidence suggests that hippocampal and caudate-putamen memory systems are unlikely candidates for amygdala modulation in this particular task, since lesions of these structures do not affect CPP acquisition for either amphetamine (e.g., Hiroi & White, 1991), or food (McDonald & White, 1993). However, the nucleus accumbens receives prominent projections from the basolateral amygdala (Kita & Kitai, 1990; Krettek & Price, 1978; McDonald, 1991; Russchen & Price, 1984), and evidence from pharmacological and lesion studies implicate the nucleus accumbens in the acquisition and expression of CPPs for various drug treatments and naturally occurring stimuli (for reviews, see Carr et al., 1989; Schechter & Calcagnotto, 1993). Moreover, the nucleus accumbens appears to interact with basolateral amygdala function in stimulus-reward learning (for reviews see Cador et al., 1989; Everitt & Robbins, 1992). Amygdalo-striatal projections are unidirectional in nature (Kita & Kitai, 1990), and this pathway has been suggested to provide the basolateral amygdala access to motor output involved in behavioral expression of stimulus-reward learning (e.g., Everitt et al., 1991). Alternatively, this pathway could conceivably allow the basolateral amygdala to exert a modulatory influence on long-term synaptic plasticity in the nucleus accumbens, a time-limited process in which behavioral expression would ultimately be independent of amygdala function. The latter suggestion is consistent with the present observation of lidocaine-induced blockade of memory consolidation, but not the expression of a food CPP, and with recent findings indicating that lesions of the basolateral amygdala block the ability of drug-associated stimuli to reinstate responding for cocaine after 21 days, but not after 43 days of drug withdrawal (Meil & See, 1997). Thus, further studies examining basolateral amygdala–nucleus accumbens interactions may aid our understanding of the modulatory role of the amygdala in consolidation of stimulus-reward memory.

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