Blockade of dopamine activity in the nucleus accumbens impairs learning extinction of conditioned fear

Orit Holtzman-Assif, Vincent Laurent, and R. Frederick Westbrook

Three experiments used rats to investigate the role of dopamine activity in learning to inhibit conditioned fear responses (freezing) in extinction. In Experiment 1, rats systemically injected with the D2 dopamine antagonist, haloperidol, froze more across multiple extinction sessions and on a drug-free retention test than control rats. In Experiment 2, rats extinguished under an intracerebroventricular (ICV) infusion of haloperidol suppressed fear responses across extinction but froze more on a subsequent drug-free retention test than control rats. In Experiment 3, rats extinguished under an infusion of haloperidol in the nucleus accumbens were impaired in suppressing fear responses across extinction and froze more on subsequent drug-free retention test than control rats. These results show that learning to inhibit fear responses in extinction requires dopamine activity in the nucleus accumbens. They were interpreted to mean that dopaminergic activity in the nucleus accumbens regulates the prediction error required for learning to inhibit fear responses in extinction.

Moreover, the omission of a predicted reward is accompanied by a suppression of firing at the time when the US normally occurred. There is also evidence from aversive conditioning procedures that DA levels in the nucleus accumbens (Acb) are increased following presentations of an unexpected footshock and that these levels are also increased by a CS which has come to signal the footshock US (Sorg and Kalivas 1991; Young et al. 1993; Guaracci and Kapp 1999; Young 2004). DA activity also appears to play a role in extinction (Pan et al. 2008). For instance, responses of midbrain DA neurons decline across repeated presentation of a CS in a manner that is correlated with the decline of conditioned responding. Moreover, many of these neurons come to show inhibitory responses at the end of extinction, consistent with the inhibitory nature of the learning produced. These findings raise the possibility that dopaminergic activity in the Acb is involved in extinction, potentially regulating some aspect of the prediction error necessary for the development of learned inhibition. However, the role played by Acb DA in learning inhibition in fear extinction is unknown.

The present experiments examined the role played by DA in learning extinction of conditioned fear responses (i.e., freezing; see Blanchard and Blanchard 1969). Seventy-one experimentally naive male Wistar rats (Gore Hill Research Laboratories, Sydney, New South Wales, Australia) were used. They were kept in a climate-controlled colony room under natural lighting. Seven days after arrival in the laboratory, rats were handled each day for 4 d. On day 1 of Experiment 1, rats were placed in conditioning chambers and were trained to fear a clicker CS (30 sec, 75 dB) that co-terminated with a footshock US (1 sec, 0.6 mA). There were four pairings of the clicker CS and the footshock US. On day 2, rats were injected i.p. with either vehicle (Groups Extinction: 0 mg/kg [n = 10] and No-extinction: 0 mg/kg [n = 8]) or one of the three doses of haloperidol (Groups Extinction: 0.05 mg/kg [n = 6], Extinction: 0.1 mg/kg [n = 7], Extinction: 1 mg/kg [n = 6], and No-extinction: 0.1 mg/kg [n = 8]) selected on the basis of previous research (Joseph et al. 2000; Shadach et al. 2000). Forty-five minutes later, rats in Groups Extinction received an extinction session in which they were placed in the conditioning chambers and presented 2 min later with the clicker CS for 5 min in the absence of the footshock.
US. Rats in the control groups (Group No-extinction) were simply handled. These procedures were repeated on days 3 and 4. On day 6, all rats were tested drug-free. Two minutes after placement in the chambers, rats were presented with the CS for 10 min in the absence of the footshock US.

Levels of freezing to the clicker CS across extinction and test are shown in Figure 1. In this and the remaining experiments, the data were analyzed by a set of planned contrasts that controlled the error rate using the Bonferroni inequality procedure (α = 0.05). Conditioning on day 1 was successful: All rats exhibited substantial levels of freezing when exposed to the clicker on day 2 (Fig. 1A). Haloperidol impaired extinction. Rats treated with the D2 antagonist froze significantly more across the first extinction session on day 2 (Fig. 1A) than vehicle-treated rats (F(1,25) = 27.24, P < 0.05). Rats that received the two highest doses of haloperidol froze at similar levels (F(1,25) = 0.4, P > 0.05), and these levels did not differ from those by rats given the lowest dose (F(1,25) = 1.4, P > 0.05). The next two extinction sessions on days 3 and 4 (Fig. 1B,C) confirmed that haloperidol impaired extinction in a dose-dependent manner. Rats injected with the drug froze significantly more across the two sessions than vehicle-treated rats (F(1,25) = 45.2, P < 0.05 and F(1,25) = 20.7, P < 0.05). Rats that received the two highest doses of haloperidol froze at similar levels (F(1,25) = 0.6, P > 0.05 and F(1,25) = 0, P > 0.05), and these levels were significantly more than those by rats given the lowest dose (Group Extinction: 0.05 mg/kg; F(1,25) = 18.6, P < 0.05 and F(1,25) = 7.5, P < 0.05). The overall levels of freezing declined from day 2 to day 4 (F(1,25) = 20.4, P < 0.05).

The impairment in extinction on day 1 could have been due to potentiation of freezing responses among haloperidol-treated rats as they froze significantly more in the pre-CS period than vehicle-treated rats (F(1,25) = 9.5, P < 0.05). However, this difference in freezing was absent on days 3 and 4 (F(1,25) = 0.9, P > 0.05 and F(1,25) = 0.1, P > 0.05). A detailed analysis of the pre-CS period on these days revealed that rats in Group Extinction 1 mg/kg froze significantly less than rats in Group Extinction 0.1 mg/kg (F(1,25) = 14.7, P < 0.05 and F(1,25) = 15.7, P < 0.05). This suggests that pre-CS freezing cannot account for the impairment in CS extinction as both doses impaired that extinction. Further, rats in Group Extinction 0.05 mg/kg froze just as much as rats that received the two higher doses of haloperidol in the pre-CS period on days 3 and 4 but nevertheless froze less to the CS than rats given the higher doses.

The test data (Fig. 1D) show that systemic injection of haloperidol had impaired long-term retention of any inhibition acquired across extinction. Rats that had not received extinction (Groups No-extinction) froze significantly more than all the other groups of rats (F(1,39) = 16, P < 0.05). The injection of haloperidol by itself was without effect since rats in Groups No-extinction showed equivalent levels of freezing whether they had been injected with saline or haloperidol but in the absence of extinction (F(1,39) = 0.1, P > 0.05). Rats in Group Extinction–Vehicle froze significantly less than rats extinguished under the drug (Groups Extinction Extinction 0.05 mg/kg, Extinction 0.1 mg/kg, and Extinction 1 mg/kg) (F(1,39) = 12.5, P < 0.05). But there were no statistically significant differences between the levels of freezing among the rats extinguished under the lowest versus the two higher doses (F(1,39) = 0.1, P > 0.05) or between rats extinguished under the two higher doses, (F(1,39) = 0.2, P > 0.05). Rats in Group Extinction 0.05 mg/kg froze significantly more than all the other groups of rats in the pre-CS period on test (F(1,39) = 10.4, P < 0.05), but there were no significant differences among the remaining groups (Fs < 4).

Experiment 1 showed that systemic haloperidol impaired the inhibition of fear responses in extinction and retention of that inhibition. Experiment 2 examined whether the impairment produced by haloperidol was observed when the drug was infused into the brain. Under anesthesia, rats were implanted with cannula in the right ventricle (AP = −0.8; ML = +1.5; DV = −4). Following one week for recovery from surgery, rats were trained to fear the clicker CS on day 1 in the manner described. On day 2, rats were infused in the ventricle with either vehicle (2 μL) (Group Extinction–Vehicle, n = 6) or haloperidol (2 μg/2 μL) (Group Extinction–Haloperidol, n = 7). Fifteen minutes later, rats received an extinction session. This consisted in placing the rats in the

Figure 1. Systemic haloperidol impairs extinction of conditioned fear. All illustrations show the mean and SEM levels of freezing. (A–C) Control rats (Group Extinction: 0 mg/kg) suppressed freezing responses across the three extinction sessions. Haloperidol impaired the suppression of freezing responses (Groups Extinction: 0.05 mg/kg, Extinction: 0.1 mg/kg, and Extinction: 1 mg/kg). (D) The retention test showed that haloperidol (Groups Extinction: 0.05 mg/kg, Extinction: 0.1 mg/kg, and Extinction: 1 mg/kg) impaired long-term retention of extinction.
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Conditioning chambers and, 2 min later, presenting the CS for 10 min in the absence of the US. On day 4, all rats were tested drug-free in the chambers. Two minutes after placement in the chambers, rats were presented with the CS for 5 min in the absence of the US.

Levels of freezing to the clicker CS across extinction and test are shown in Figure 2. Conditioning on day 1 was successful as all rats exhibited substantial levels of freezing when presented with the clicker on day 2 (Fig. 2A). Haloperidol did not affect fear responses across extinction. Rats infused with haloperidol into the ventricle froze just as much as did rats infused with vehicle ($F_{(1,11)} = 0.3, P > 0.05$). Overall levels of freezing significantly decreased across the extinction session ($F_{(1,11)} = 74.9, P < 0.05$).

The test data (Fig. 2B) show that intracerebroventricular (ICV) infusion of haloperidol had impaired the retention of the inhibition produced by the CS-alone exposure. Rats that had been extinguished to the CS under haloperidol froze significantly more than rats extinguished under vehicle ($F_{(1,11)} = 8.4, P < 0.05$). There were no statistically significant differences between the groups in the period prior to the presentation of the CS on test ($F_{(1,11)} = 0.8, P > 0.05$).

Experiment 2 showed that an ICV infusion of the DA antagonist impaired retention of inhibitory learning. Experiment 3 examined whether the Acb constituted one of the sites at which the drug had acted to impair retention. The procedure was identical to that used in Experiment 2, except that rats were infused bilaterally in the Acb (AP $= +1.4$; ML $= 1.5$; DV $= -7$) with either vehicle (0.25 $\mu$L per side) or haloperidol (0.25 $\mu$g/0.25 $\mu$L per side) before the extinction session on day 2.

Levels of freezing to the clicker CS across extinction and on test are shown in Figure 3. Conditioning on day 1 was successful as all rats froze when re-exposed to the CS on day 2 (Fig. 3B).

Haloperidol impaired the depressive effects of extinction. Rats in Group Extinction–Haloperidol ($n = 6$) froze significantly more than rats in Group Extinction–Vehicle ($n = 7$; $F_{(1,11)} = 5.1, P < 0.05$). Overall levels of freezing significantly decreased across the extinction session ($F_{(1,11)} = 5.3, P < 0.05$). Haloperidol-treated rats froze at similar levels as did rats treated with vehicle in the period prior to presentation of the clicker on day 2 ($F_{(1,11)} = 2.9, P > 0.05$).

The retention test data (Fig. 3C) show that intra-accumbal infusion of haloperidol impaired retention of inhibition. Rats extinguished under the drug froze significantly more on the retention test than rats extinguished under vehicle ($F_{(1,11)} = 17.3, P < 0.05$). There were no statistically significant differences between the groups in the period prior to presentation of the clicker CS on test ($F_{(1,11)} = 0.9, P > 0.05$).

The present series of experiments examined the role of dopaminergic activity in learning to inhibit conditioned fear responses in extinction. In Experiment 1, rats that received systemic injection of the D2 DA antagonist, haloperidol, before extinction showed more fear responses across a subsequent drug-free test than rats injected with vehicle. The drug impaired retention of inhibition across each of the three doses used. The drug per se did not elevate the test levels of freezing since control rats injected with the drug but not subjected to extinction did not differ across test from vehicle-injected rats not subjected to extinction. However, haloperidol-treated rats showed more freezing across each of the extinction sessions than control rats, suggesting that the drug had acted to either potentiate freezing or block the development of its inhibition. Alternatively, rats extinguished under the drug may have shown substantial levels of freezing on test not because inhibition was impaired but because of generalization decrement between the drug cues present across extinction and their absence on test, that is, because of state dependency of the learning (Overton 1985). However, the D2 DA antagonist was administered before each of the three extinction sessions conducted across consecutive days. Thus, the second and third sessions can be viewed as a retention test in the presence of haloperidol for the inhibition learned the previous day. These tests also showed that haloperidol impaired retention of inhibition. Further, the next two experiments clearly rule out a critical role for either a potentiation of fear responses or state dependency in the impairment of inhibition produced by haloperidol. In Experiment 2, rats infused before extinction with the D2 DA antagonist in the right ventricle froze more than control rats across the drug-free test given the following day. Importantly, those haloperidol-treated rats froze just as much as control rats across extinction, suggesting that DA activity is involved in consolidation of inhibition. Experiment 3 extended these findings by showing that dopaminergic activity in the nucleus accumbens is particularly important for extinction. Rats that received Acb infusion of haloperidol exhibited more freezing across extinction than control rats and also more freezing across the subsequent drug-free test. Thus, accumbal DA appears to be critical for learning to inhibit conditioned fear responses in extinction and for retaining that inhibition.

The finding that accumbal DA activity regulates the development and retention of the fear inhibition produced by extinction is consistent with its proposed role in predictive learning (Schultz and Dickinson 2000). Contemporary learning theories use error correction mechanisms of one sort or another to explain a range of acquisition phenomena. Such theories also hold that the omission of the predicted US constitutes the error which drives the inhibitory association that comes to suppress fear responding in extinction (Dickinson 1980; Rescorla 1988). The present experiments suggest that changes in the levels of DA in the nucleus accumbens are involved in coding for this error or in transmitting the error signal into other brain regions. This suggestion is consistent with a recent study investigating the role of...
midbrain DA in extinction of appetitive conditioning (Pan et al. 2008). It revealed that the activity of these neurons declines across repeated presentation of the CS in a manner that is correlated with the decline of the conditioned response. Moreover, many of these neurons came to show inhibitory responses at the end of extinction, consistent with the inhibitory learning assumed to underlie the suppression of responding across extinction. Although these results were obtained in DA neurons located in the VTA, our results provide evidence that accumbal DA also modulates predictive learning in extinction. For example, the local infusion of haloperidol may have prevented the development of inhibitory responses in accumbal DA neurons and thereby impaired extinction of conditioned fear responses.

Regardless of these suggestions concerning its role in prediction error, the present experiments clearly show that DA in the nucleus accumbens is critical for learning and retention of the inhibition produced by fear extinction. However, a currently popular neural model (Quirk and Mueller 2008) does not incorporate a role for the nucleus accumbens in learning extinction. This model attributes extinction to interactions among several structures, but most notably the BLA and the mPFC. Specifically, acquisition and consolidation of the inhibition formed in extinction requires activity in the BLA (Lin et al. 2003; Herry et al. 2006; Kim et al. 2007; Sotres-Bayon et al. 2007, 2009; Laurent and Westbrook 2008; Laurent et al. 2008). In contrast, the infralimbic (IL) subregion of the mPFC is required for consolidation and retrieval of that inhibition (Quirk et al. 2000; Milad and Quirk 2002; Santini et al. 2004; Hugues et al. 2006; Sierra-Mercado et al. 2006; Burgos-Robles et al. 2007; Laurent and Westbrook 2008, 2009; Sotres-Bayon et al. 2009). Further, the IL is held to suppress fear responses to an extinguished CS through projections to a network of inhibitory interneurons located between the BLA and the CeA (Pare and Smith 1993; McDonald 1998; Royer et al. 1999). Lesioning these interneurons disrupts the expression of extinction (Likhtik et al. 2008), and electrical stimulation of the IL reduces fear responses as well as the activity of CeA neurons that are responsible for eliciting those responses (Vidal-Gonzalez et al. 2006). This model is consistent with much of the available data on extinction, but needs to be extended to include the error correction mechanisms that regulate inhibitory learning in fear extinction. Our results indicate that such mechanisms may involve changes in accumbal DA. The Acb receives projections from both the BLA and the mPFC and is also a major target for DA neurons originating in the VTA (Groenewegen et al. 1990, 1999; Wright et al. 1996). This connectivity suggests that the Acb receives information about the CS from both cortical regions and the BLA, as well as information about the outcome of the CS presentation from the VTA. Accordingly, the Acb may function as a device that holds information about the CS that is then processed differently depending on the error signal provided by DA activation. It is noteworthy that the Acb sends indirect projections to both the BLA, where the initial inhibition is encoded, and the IL cortex, where it is stored (Groenewegen et al. 1990, 1999; Wright et al. 1996). Thus, the release of DA into the Acb may regulate interactions between the BLA and the mPFC that are necessary for learning inhibition in extinction.

References


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