Antagonism of lateral amygdala alpha1-adrenergic receptors facilitates fear conditioning and long-term potentiation

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Norepinephrine receptors have been studied in emotion, memory, and attention. However, the role of alpha1-adrenergic receptors in fear conditioning, a major model of emotional learning, is poorly understood. We examined the effect of terazosin, an alpha1-adrenergic receptor antagonist, on cued fear conditioning. Systemic or intra-lateral amygdala terazosin delivered before conditioning enhanced short- and long-term memory. Terazosin delivered after conditioning did not affect consolidation. In vitro, terazosin impaired lateral amygdala inhibitory postsynaptic currents leading to facilitation of excitatory postsynaptic currents and long-term potentiation. Since alpha blockers are prescribed for hypertension and post-traumatic stress disorder, these results may have important clinical implications.

Although norepinephrine (NE) has been widely studied as an important modulator of memory and emotion, comparatively little is known about the role of NE in amygdala-dependent Pavlovian fear conditioning, a major model for understanding the neural basis of fear learning and memory. In fear conditioning, an emotionally neutral conditioned stimulus (CS; i.e., tone) is temporally paired with an aversive unconditioned stimulus (US; i.e., footshock). After very few pairings, a lasting, robust CS–US association is acquired, and the CS elicits stereotypical Pavlovian fear responses.

We hypothesized that alpha1 blockers would facilitate fear learning, possibly by impairing LA inhibition and facilitating LA LTP. To test this hypothesis, we injected rats with terazosin, a selective alpha1-adrenergic receptor antagonist, systemically or directly into LA before or after fear conditioning. We examined in vitro the effect of terazosin on LA pyramidal cell inhibitory postsynaptic currents (IPSCs) and excitatory postsynaptic currents (EPSCs) in response to fiber stimulation of the thalamic CS input pathway to LA, as well as the effect of terazosin on LA LTP in this same pathway. We found that intra-LA terazosin facilitated fear conditioning when injected before but not after training. We also found that terazosin impaired IPSCs in LA pyramidal cells, leading to facilitated EPSCs and LTP.

Behavioral experiments were conducted on adult, male Sprague–Dawley rats (Hilltop Laboratory Animals) weighing approximately 300 g upon arrival. Rats were individually housed, maintained on a 12/12 h light/dark schedule, and allowed free access to food and water. Testing was conducted during the light phase. All procedures and experiments were approved by NYU’s Animal Care and Use Committee.

For systemic injections, terazosin (20 mg/kg; Sigma) was dissolved in aCSF and infused into the LA at 0.1 μL/min 30 min prior to or immediately after fear conditioning. Bilateral guide cannulae (22 gauge; Plastics One) aimed at LA (~3.3 mm anterior, 5.2 mm lateral, ~7 mm dorsal to bregma) were surgically implanted as previously described (Sotres-Bayon et al. 2009). Rats were given at least 7 d to recover.
from surgery before testing. For infusions, dummy cannulae were removed, and infusion cannulae (28 gauge, +1 mm beyond guides) were inserted into guides. Infusion cannulae were connected to a 1.0 µL Hamilton syringe via polyethylene tubing. Infusion rate was controlled by a pump (PHD22; 2000; Harvard Apparatus), and infusion cannulae were left in place for an additional 60 sec to allow diffusion of the solution away from the cannula tip, then dummy cannulae were replaced. Upon completion of the experiment, rats were euthanized, brains removed, and cannulae placements verified histologically as previously described (Sotres-Bayon et al. 2009).

Two contexts (A and B) were used for all testing as previously described (Schiller et al. 2008). The grid floors in Context B were covered with black Plexiglas inserts to reduce generalization. NO odors were used and chambers were cleaned between sessions. CSs were 30 sec, 5 kHz, 80 dB tones, and USs were 1 sec, 0.8 mA scrambled electric footshocks. Experiments consisted of two phases separated by 48 h: (1) fear conditioning in Context A and (2) long-term memory (LTM) test in Context B. On Day 1, rats were placed in Context A, allowed 5 min to acclimate, and then received three CS-US pairings separated by variable 5 min ITIs. On Day 3, rats were placed in Context B and allowed 5 min to acclimate before receiving one CS-alone presentation.

The index of fear in behavioral experiments was “freezing,” the absence of all non-respiratory movement (Blanchard and Blanchard 1971; Fanselow 1980). Following testing, freezing was manually scored from DVDs by a scorer blind to group specifications. Graphs represent group means ± SEM. Statistical analysis was conducted with GraphPad Prism.

Whole-cell electrophysiological recordings were obtained from LA pyramidal cells using in vitro coronal slices from rats aged P21–P30 d as described in Cunha et al. (2010). Terazosin was bath-applied for 10 min to achieve stable responses before testing. The cells were voltage-clamped using an Axopatch 200B amplifier at −35 mV for recording EPSCs and IPSCs. Synaptic responses were evoked with sharpened tungsten bipolar stimulating electrodes. Internal capsule fibers containing thalamic afferents were stimulated for paired-pulse facilitation (PPF) (ISI = 50 msec; 0.1 Hz) using a photoelectric stimulus isolation unit with a constant current output. Cells were rejected if access resistance (8–26 MΩ) changed more than 15%. Signals were filtered at 2 kHz and digitized (Digitida 1440 A; Axon Instruments), and peak amplitude, 10%–90% rise time, and IPSC decay time constants were analyzed offline using pCLAMP10.2 software (Axon Instruments).

Brain slices for LTP experiments were prepared from rats aged 3–5 wk as in Johnson et al. (2008) and maintained on an interface chamber at 31 °C. Glass recording electrodes (filled with aCSF, 5 MΩ resistance) were guided to LA neurons. Bipolar stainless steel stimulating electrodes (75 kΩ) were positioned medial to LA in internal capsule fibers. Orthodromic synaptic potentials were evoked via an isolated current generator (Digitimer; 100 μsec pulses of 0.3–0.7 mA). Evoked field potentials were recorded with an Axoclamp 2B amplifier and Axon WCP software (Axon Instruments). Data were analyzed offline using WCP PeakFit (Axon Instruments). LTP was measured as a change in evoked field potential amplitude.

Baseline responses were monitored at 0.05 Hz for 30 min with a stimulus intensity of 40%–50% of maximum field potential before LTP induction. Terazosin (10 µM) was superfused for 15 min, and then LTP was elicited by three tetanus trains (100 Hz × 1 sec, 3 min ITI) with the same intensity and pulse duration as the baseline stimuli. In one experiment, picrotoxin (PTX; 75 µM) was present in the perfusion solution to block fast GABAergic signaling.

**Figure 1.** Pre-training terazosin facilitates conditioned fear learning when injected systemically or directly into LA. (A) Pre-CS- and CS-elicited freezing behavior during fear conditioning (left) and during a test CS 48 h after fear conditioning (right). Terazosin (n = 10) or saline (n = 9) was injected i.p. 30 min before conditioning. (B) Identical experiment except terazosin (n = 8) or aCSF (n = 7) was microinfused into LA 30 min prior to fear conditioning. (C) Identical experiment except terazosin (n = 6) or aCSF (n = 6) was microinfused into LA immediately after fear conditioning. Arrows indicate when drug treatment occurred. (*) P < 0.05 versus control.

**Terazosin facilitates acquisition, but not consolidation, of fear conditioning**

We first examined the effect of terazosin, injected systemically 30 min before training, on fear learning and memory (Fig. 1A). Learning and short-term memory effects were assessed as differences in freezing during the second and third CSs of the three trial acquisition sessions. LTM effects were assessed as differences in freezing during the drug-free expression test CS 48 h after training. Although systemic terazosin caused sedation and reduced locomotion in some rats, it also increased CS-elicited freezing relative to controls, suggesting a learning facilitation. Freezing was distinguished from sedation by differences in posture and whisker tension. Two-way ANOVA of freezing scores during training revealed a significant effect of the group × trial interaction (F2,340 = 3.06, P < 0.05), and post-hoc Bonferroni tests showed a significant difference in freezing for the second (P < 0.01) and third (P < 0.01) CSs. Freezing was significantly elevated in terazosin-treated rats during the drug-free LTM test (t(1,17) = 3.09, P < 0.01).
We next infused terazosin directly into LA prior to fear conditioning to localize the drug effect and minimize concerns about nonspecific effects (sedation) associated with systemic injections. Intra-LA terazosin facilitated fear learning and LTM relative to control rats (Fig. 1B). Two-way ANOVA of freezing scores during training revealed a significant group × trial interaction effect ($F_{(29,290)} = 4.73, P < 0.05$), and post-hoc Bonferroni tests showed a significant difference in freezing for the second CS ($P < 0.01$). For the LTM test, intra-LA terazosin-treated rats froze more than control rats ($t_{(1,9)} = 2.556, P < 0.05$).

To differentiate between potential learning versus memory consolidation effects, we infused terazosin into LA immediately after fear conditioning in separate rats (Fig. 1C). Freezing was similar for terazosin- or vehicle-treated rats at the LTM test ($t_{(1,8)} = 0.37$), suggesting that terazosin facilitates learning, but not consolidation, of fear conditioning.

**Terazosin facilitates LTP in LA, most likely by impairing local inhibition**

We first evaluated the effect of terazosin on thalamo-LA EPSCs and IPSCs. Synaptic current traces of an LA pyramidal cell ($n = 11$) under applied terazosin and control conditions are shown in Figure 2A. An application of 10 μM terazosin significantly decreased IPSCs 80.7% ± 17.2% ($p'V = 0.013$) and led to PPF of EPSCs 64.3% ± 14.5% ($p'V = 0.006$).

We next evaluated the effect of terazosin on thalamo-LA LTP induction with and without PTX. Without PTX, terazosin facilitated LTP (Fig. 2B). Two-way ANOVA of evoked responses post-LTP-induction showed a significant effect of the group × time interaction ($F_{(29,319)} = 5.97, P < 0.01$) and time ($F_{(29,319)} = 1.71, P < 0.05$). With PTX present (Fig. 2C), terazosin slightly impaired LTP, again showing a significant effect of interaction ($F_{(29,290)} = 1.63, P < 0.05$) and time ($F_{(29,290)} = 1.82, P < 0.01$). We also conducted identical LTP experiments in the cortex→LA pathway (stimulating external capsule), and terazosin failed to facilitate LTP without PTX and produced a modest impairment with PTX (data not shown). This null effect in the cortical pathway highlights the importance of the thalamic pathway in fear conditioning and suggests that α1 receptors mainly constrain thalamo-LA plasticity underlying memory formation.

Noradrenergic modulation of memory and anxiety has been studied for decades (Gray 1978; Davis et al. 1979; Redmond and Huang 1979). However, less is known about the role of NE in Pavlovian fear conditioning, a major model for understanding the biological basis of memory and generation of normal and pathological fear. Of the major noradrenergic receptor classes, α1 receptors have received less attention in studies of memory and fear than both the beta and alpha2 receptors. Because α1 receptors are strongly expressed in LA, a key region of plasticity in fear conditioning, we chose to examine the effects of terazosin on amygdala-dependent Pavlovian fear conditioning and its underlying synaptic processes. We found that systemic and intra-LA terazosin significantly enhanced fear learning without affecting memory consolidation. Terazosin also facilitated synaptic plasticity in an important LA fear conditioning pathway, most likely by impairing local GABAergic inhibition. These findings provide important insight into the neuromodulatory mechanisms affecting fear learning and may have important clinical implications.

We initially investigated the role of α1 receptors in fear conditioning using the α1 blocker prazosin injected systemically before training (Cain et al. 2006). Prazosin enhanced CS-elicited freezing within the training session and during the drug-free LTM test, suggesting that α1 receptors contribute to a constraining process in fear conditioning. We switched to terazosin in these studies because it is approximately 25 times more soluble than prazosin and more selective for the α1 receptors, although slightly less potent (Achari and Laddu 1992). Thus, terazosin was better for investigating α1 contributions to brain processes important for fear conditioning. Like prazosin, we found that systemic terazosin facilitated CS-elicited freezing both within the training session and the drug-free LTM test (Fig. 1A). It is important to note that terazosin at the dose tested (20 mg/kg) did appear to induce sedation in some rats, possibly confounding within-session freezing scores.

To localize terazosin effects to a brain site and circumvent concerns about nonspecific systemic drug effects, we infused terazosin directly into LA before fear conditioning. Again, terazosin enhanced CS-elicited freezing during...
training and during the drug-free LTM test (Fig. 1B). There were no signs of sedation with intra-amygdala infusions. The within-session enhancement of freezing suggests that LA alpha1 receptors normally act to constrain learning-related processes. NE, however, has been strongly implicated in consolidation processes with other fear-related behavioral paradigms such as inhibitory avoidance (Ferry and McGaugh 2008). Thus, we directly examined the contribution of LA alpha1 receptors to consolidation of fear conditioning by infusing terazosin immediately after training (Fig. 1C). We found no significant effect on CS-elicited freezing during the LTM test. Together, our findings implicate LA alpha1 receptors in a process that constrains acquisition, but not consolidation, of Pavlovian fear conditioning.

Although alpha1 receptors have never been directly examined in a cue fear conditioning procedure, amygdala-alpha1 receptor activity has been implicated in consolidation of inhibitory avoidance (McGaugh 2004). We do not believe that our findings contradict the avoidance findings. While the two procedures are related, they represent different forms of learning that depend on different brain regions. Fear conditioning is a Pavlovian procedure in which animals learn associations between environmental stimuli but have no control over their delivery. Inhibitory avoidance has an instrumental component—the animal withholds responses that lead to shock delivery. Additionally, whereas Pavlovian conditioning critically depends on LA for learning and memory, inhibitory avoidance appears to depend on LA mainly for modulation of consolidation processes taking place outside of the amygdala (McGaugh et al. 2002). It remains possible that LA alpha1 receptors can contribute both to acquisition of Pavlovian fear conditioning and consolidation of inhibitory avoidance. Future studies will be necessary to disentangle the exact contribution of LA alpha1 receptors to these two fear-related learning paradigms. One possibility is that alpha1 receptors on different neuron types (excitatory vs. inhibitory) or different populations of excitatory neurons explains the differential effects of alpha1 blockers in the two paradigms. This dissociation of noradrenergic contributions to fear conditioning and inhibitory avoidance is also seen with manipulations of beta-adrenergic receptors, where the antagonist propranolol blocks consolidation of inhibitory avoidance (McGaugh et al. 2002) but not fear conditioning (Lee et al. 2001; Debiec and LeDoux 2004; Bush et al. 2010).

Two ways NE may modulate fear acquisition have been identified. First, NE may amplify Hebbian mechanisms within pyramidal neurons that acquire plastic changes underlying conditioning (Braga et al. 2004) and terazosin, appear to disinhibit plasticity in LA fear conditioning. Consistent with our findings, alpha1 receptor activity has been previously reported to participate in feed-forward inhibition in LA (Braga et al. 2004). Our PTX experiment suggests that alpha1 receptors on pyramidal cells may actually enhance Hebbian LTP processes, but under normal conditions this contribution appears to be overshadowed by their contribution to inhibitory gating.

Our results suggest that alpha1-adrenergic receptor activity in LA contributes to feed-forward inhibition and constrains fear learning. Alpha1-adrenergic receptor antagonists, like prazosin and terazosin, appear to disinhibit plasticity in LA fear conditioning pathways even when given systemically. These results demonstrate another mechanism whereby NE neuromodulation can powerfully affect learning and fear. These same alpha1 blockers are commonly prescribed to combat both hypertension and nightmares associated with PTSD. Our findings suggest that clinicians should be especially aware of patients subjected to trauma while taking alpha1 blockers, as this may lead to stronger fear learning and possible exacerbation of existing fear-related disorders.

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